### **Extended Supplementary Information for**

### **Cell-Size Control and Homeostasis in Bacteria**

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### I. EXPERIMENTAL MATERIALS AND METHODS

### A. Strain and growth conditions

Strains: For physiological study, it is important to use a prototrophic strain. For  $E.\ coli$  experiments, we chose  $E.\ coli$  K12 NCM3722 constructed, sequenced, and extensively tested by Sydney Kustu's lab [1]. We used SJ202, a nonmotile derivative of NCM3722 ( $\Delta$ motA). For  $B.\ subtilis$  experiments, we chose a strain in the 3610 background with Coml (Q12L) mutation to allow competence. We used a derivative with reduced motility and biofilm formation by deleting epsH and a flagellin protein hag, respectively.

Growth media: E. coli growth experiments were performed in seven different nutrient conditions. The average generation time in these conditions evenly spans from 17.1 to 51.4 minutes at 37°C. The growth medium is based on MOPS, developed by Fred Neidhardt [2] and is commercially available from Teknova (Teknova.com). B. subtilis growth experiments performed in four different growth conditions with average doubling times between 16.9 and 38.9 minutes. The details of the media used for E. coli and B. subtilis are listed in the Tables SII & SI. Prior to growth of the cells in the microfluidics device, all cultures were grown in a 37°C water bath shaker, shaking at 240 rpm.

Sample preparations: Cell culture for each experiment was carried out in three steps: seed culture, pre-culture and experimental culture, all in the growth medium of the corresponding experiment. The culture volume was 3 ml in 950 mm  $\times$  150 mm culture tubes to ensure aeration. For seed culture, cells were inoculated into growth media from -80 glycerol stock, cultured at 37°C with shaking overnight. The culture was then backdiluted 1000-fold in the identical growth medium and cultured in 37°C water bath shaker (pre-culture) until it reaches exponential phase.

The pre-culture was then diluted 1000-fold in the identical growth medium and cultured in 37°C water bath shaker (experimental culture). After a few cell doublings, the experimental culture was centrifuged, and concentrated 20 folds in the identical media containing 50mg/ml of bovine serum albumin (BSA), to reduce nonspecific adsorption of cells to glass surface in the microfluidic device. The concentrated culture was injected in the microfluidics device for single cell imaging. Fresh pre-warmed media was pumped in the device immediately after loading of the cells in the growth channels.

Media name	Buffer	Carbon source	Supplement	Mass doubling
(as used in the text)		(v/w) concentration		time at $37^{\circ}$ C
glycerol	MOPS modified buffer	Glycerol 0.4%	-	51.4
sorbitol	MOPS modified buffer	Sorbitol 0.2%	_	50.9
glucose	MOPS modified buffer	Glucose $0.2\%$	=	37.7
glucose + 6 a.a.	MOPS modified buffer	Glucose $0.2\%$	table I.b	30.1
glucose $+ 12$ a.a.	MOPS modified buffer	Glucose $0.2\%$	tables I.b and I.c	26.6
Synthetic rich	MOPS modified buffer	Glucose $0.2\%$	table I.d	22.5
Tryptic soy broth (TSB)	Tryptone 1.7%, Soytor			17.1
Tryptic soy broth (15b)	glucose 0.25%, and dip	otassium phosphate (	0.25%	17.1

Table 1	I a·	MOPS	modified	buffer

Components	Concentration
MOPS (MW 209.3)	40mM
Tricine (MW 179.2)	$4.0 \mathrm{mM}$
Iron Sulfate Stock	$0.01 \mathrm{mM}$
Ammonium Chloride	$9.5 \mathrm{mM}$
Potassium Sulfate	$0.276 \mathrm{mM}$
Calcium Chloride	$0.0005 \mathrm{mM}$
Magnesium Chloride	$0.525 \mathrm{mM}$
Sodium Chloride	$50 \mathrm{mM}$
Ammonium Molybdate	$3 \times 10^{-9} \text{ M}$
Boric Acid	$4 \times 10^{-7} \text{ M}$
Cobalt Chloride	$3 \times 10^{-8} \text{ M}$
Cupric Sulfate	$10^{-8} \text{ M}$
Manganese Chloride	$8 \times 10^{-8} \text{ M}$
Zinc Sulfate	$10^{-8} \text{ M}$
Potassium Phosphate Dibasic	$1.32 \mathrm{mM}$

Table I.b: supplements for glucose+6 a.a. and +12 a.a.

Components	Concentration ( $\mu g/ml$ )
L-Methionine	5
L-Histidine	5
L-Arginine	5
L-Proline	5
L-Threonine	5
L-Tryptophan	5

Table I.c: supplements for glucose+12 a.a.

Components	Concentration ( $\mu$ g/ml)
L-Serine	5
L-Leucine	5
L-Tyrosine	5
L-Alanine	5
L-Asparagine	5
L-Aspartic Acid	0.25

Table I.d: supplements for synthetic rich media

Components	Concentration
L-Alanine	$0.8 \mathrm{mM}$
L-Arginine	$5.2 \mathrm{mM}$
L-Asparigine	$0.4 \mathrm{mM}$
L-Aspartic Acid, Potassium Salt	$0.4 \mathrm{mM}$
L-Glutamic Acid, Potassium Salt	$0.66 \mathrm{mM}$
L-Glutamine	$0.6 \mathrm{mM}$
L-Glycine	$0.8 \mathrm{mM}$
L-Histidine HCl H <sub>2</sub> O	$0.2 \mathrm{mM}$
L-Isoleucine	$0.4 \mathrm{mM}$
L-Proline	$0.4 \mathrm{mM}$
L-Serine	$10 \mathrm{mM}$
L-Threonine	$0.4 \mathrm{mM}$
L-Tryptophan	$0.1 \mathrm{mM}$
L-Valine	$0.6 \mathrm{mM}$
L-Leucine	$0.8 \mathrm{mM}$
L-Lysine	$0.4 \mathrm{mM}$
L-Methionine	$0.2 \mathrm{mM}$
L-Phenylalinine	$0.4 \mathrm{mM}$
L-Cysteine HCl	$0.1 \mathrm{mM}$
L-Tyrosine	$0.2 \mathrm{mM}$
Thiamine	$0.01 \mathrm{mM}$
Calcium Pantothenate	$0.01 \mathrm{mM}$
para-Amino Benzoic Acid	$0.01 \mathrm{mM}$
para-Hydroxy benzoic Acid	$0.01 \mathrm{mM}$
di Hydroxy Benzoic Acid	$0.01 \mathrm{mM}$
Potassium Hydroxide	$1.5 \mathrm{mM}$
Adenine	$0.2 \mathrm{mM}$
Cytosine	$0.2 \mathrm{mM}$
Uracil	$0.2 \mathrm{mM}$
Guanine	$0.2 \mathrm{mM}$
	ļ

Table SI List of growth media, carbon sources and the supplements that we used for E. coli in this study.

Media name	Buffer	Carbon source	Supplement	Mass doubling
(as used in the text)		(v/w) concentration		time at $37^{\circ}$ C
S7 <sub>50</sub> glycerol	tables II.a and II.c	Glycerol 1%	table II.b	35.6
$S7_{50}$ glucose	tables II.a and II.c	Glucose 1%	table II.b	38.9
$_{ m LB}$	Tryptone 1%, Yeas	t extract 0.5% and N	aCl 1%	17.8
Truntia gove broth (TCD)	Tryptone 1.7%, So	ytone 0.3%, sodium	chloride $0.5\%$ ,	16.9
Tryptic soy broth (TSB)	glucose $0.25\%$ , and	dipotassium phospha	te $0.25\%$	10.9

Table II.a: $S7_{50}$ s	salts			Table II.c: S7 <sub>50</sub>	0 metals
Components		Concentration		Components	Concentration
MOPS buffer		$50 \mathrm{mM}$	•	$MgCl_2$	2mM
Ammonium sulfat	te $(NH_4)_2SO_4$	$1 \mathrm{mM}$		$CaCl_2$	$700\mu\mathrm{M}$
Potassium phosph	hate monobasic KH <sub>2</sub> PO <sub>4</sub>	5 mM		$MnCl_2$	$50\mu\mathrm{M}$
				$ZnCl_2$	$1\mu\mathrm{M}$
Table II.b: supple	emets			FeCl <sub>3</sub>	$5\mu\mathrm{M}$
Components   Con				${\bf Thiamine\text{-}HCl}$	$1 \mathrm{mM}$
glutamate 6.8n				HCl	$20\mu\mathrm{M}$
Na-Citrate 250	uM				
FeCl <sub>3</sub> 250 <sub>1</sub>	uM				

Table SII List of growth media, carbon sources and the supplements that we used for B. subtilis in this study.



**Figure S1.** A view of the environmental chamber. Equipment is fully housed in the 37°C environmental chamber in order to avoid temperature fluctuations.

### B. Temperature control

All experimental steps - from inoculation to imaging - were performed at  $37 \pm 0.1^{\circ}$ C. To this end, all equipment was stationed in a 5' x 7' environmental chamber (Figures S1 and S2) to eliminate any side effect of temperature fluctuations on the cell growth and physiology. Within the chamber, the temperature distribution was homogeneous with forced air circulation within  $\pm 0.1^{\circ}$ C, and constantly monitored at multiple locations.

### C. Microfluidics

To monitor single cell growth, we employed the microfluidic mother machine that we developed to study cell growth and the cell cycle [3]. Master molds, from each of which many PDMS microfluidic devices were cast, were fabricated using standard nanofabrication techniques (detailed protocols are available in "The Mother Machine Handbook" from the web site of Jun lab at http://jun.ucsd.edu and a video at http://www.youtube.com/watch?v=RGfb9XU5Oow). PDMS was prepared from a Sylgard 184 Silicone Elastomer kit: polymer base and curing agent were mixed in a 10 to 1 ratio, air bubbles were purged from the mixture in a vacuum chamber, the degassed mixture was poured over the master, and the devices were cured about 24 hours at 65°C. Cured PDMS has a consistency like rubber; devices were peeled from the master mold. Devices were treated with pentane and then acetone to remove residual uncured



Figure S2. A panoramic view of the interior of the environmental chamber showing the microscope and the other facilities that we employed for the growth and the imaging of live cells.

polymer from the PDMS matrix.

The inlet and outlet of the microfluidic device were bolstered with a thicker layer of PDMS. To bond the PDMS layers, the surfaces were exposed to oxygen plasma for 15 seconds at 30 watts in a Harrick Plasma system. Oxygen plasma makes exposed PDMS and glass reactive, so that covalent bonds form between surfaces brought into contact with one another. The seal between PDMS surfaces was established for 10 minutes at 65 °C.

The PDMS mother machine device was sealed to a Willco-dish glass bottom dish (Willco Wells) with no.1.5 glass thickness (170  $\mu$ m). Before use, the glass surfaces were wiped with ethanol and then immediately with water, to remove residual ethanol. As before, the PDMS and glass surfaces were exposed to oxygen plasma for 15 seconds. Then the PDMS microfluidic device was sealed to the cover slip and the bonds were allowed to set for 10 minutes at 65 °C. After constructing the device, the channel walls were incubated with 50 mg/ml BSA (Sigma) for surface passivation. The growth medium also contained 0.5 mg/ml BSA in the background.

Because the *E. coli* cell width changes with respect to the growth condition, each growth condition required a unique mother machine device optimized for the experiment.

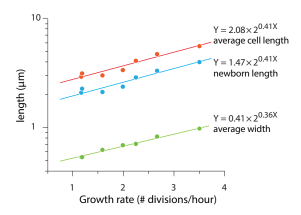
### D. Microscopy and image acquisition

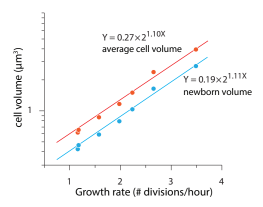
An inverted microscope (Nikon Ti-E) equipped with Perfect Focus (PFS 3), a 100x oil immersion objective lens (NA 1.45), white LED transmission light (TLED, Sutter Instruments, 400-700nm) and Andor NEO sCMOS camera was used for phase contrast imaging. The illumination condition was 50ms exposure with illumination intensity set at 10% of the max TLED intensity. The frequency of the time lapse imaging was chosen such that about 20 or more images were taken per generation time.

### E. Image analysis

We developed custom high-throughput image analysis software optimized for our growth experiments in the mother machine. The procedure includes processing of phase contrast images to eliminate background patterns from the PDMS device. The overall steps are as follows:

- Step 1: Subtract the image of an empty channel from the channels containing cells. The purpose of this step is to eliminate
  patterns from the PDMS device.
- Step 2: Find the mid-cell axis that runs through the long axis of the cells.
- Step 3: Project the brightness profile of the cells along their mid-cell axis.
- Step 4: From the brightness profile, find the cell poles and the length of the cells.





**Figure S3.** The behavior of *E. coli* width, length and volume as a function of growth rate (divisions/hour). (left) Width, length at birth and average length of the cells as a function of the growth rate. (right) Newborn cell volume, and average cell volume as a function of the growth rate. Both figures are on semi-log panels.

### II. FULL EXPERIMENTAL DATA

### A. Growth law: variation of cell volume, length and width as a function of growth rate

The growth law, based on the two classic works [4; 5], established an exponential relationship between the average cell size and the growth rate (number of divisions per hour). In our study, using single cell data, we went a step beyond and tested the growth law at the single cell level. Figure 1C of the main text confirms the exponential relationship between average volume (or, equivalently, the volume of newborn cells) and the growth rate of *E. coli*, consistent with the classic literature.

At the single cell level, we measured both the width and the length of the cell and the resulting data is shown in Figure S3. Importantly, *E. coli* cells maintained a constant width during elongation. This constancy can be seen in Figure S4A, where the cell width varies less than 1% between birth and division (by linear fit; see also Table SII). Furthermore, the width of the cell was independent of its length (Figure S4B).

Because the cell width was constant for each growth condition, we were able to calculate the volume of the cell assuming the geometry of the cell is cylindrical with two hemispherical caps. The resulting values are reported as a function of the growth rate.

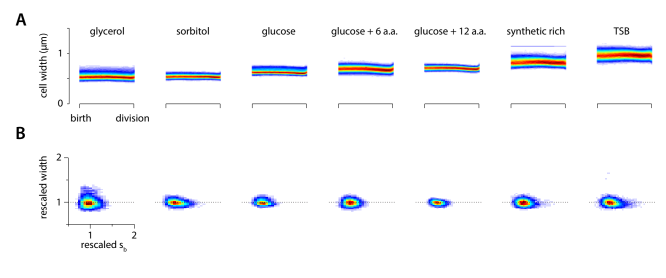
The growth law between the average newborn cell volume and the growth rate raises a question whether the newborn cell length and width also show an exponential dependence on the growth rate. Figure S3 suggests that may indeed be the case.

### B. Correlation of parameters with respect to the newborn size

In Figure 1D of the main text we presented the correlation between a set of parameters of interest, viz., the rescaled elongation rate, the generation time and the size of the cell at division, as a function of the rescaled newborn size. Here, we present data separately by showing rescaled scatter plots for the various growth conditions in Figure S5. The trend is manifestly similar for all the growth conditions.

### C. Constancy of the added size with respect to the size of the cell at birth

As a quantitative test of the  $\Delta$  model, we analyzed the distribution of the added size at division,  $\Delta_d \equiv s_d - s_b$ , where  $s_d$  and the  $s_b$  are the size of the cell at division and birth, respectively. In the SI, we shall use for size increments at division the notation  $\Delta_d$ , with the index meant to distinguish the value at division from the value during elongation that we shall need later. Figure 2 in the main text shows the distributions of the added size,  $P(\Delta_d|s_b)$ , conditional to  $s_b$ , which were observed to be independent of  $s_b$ . Here, we make a direct comparison with the conditional distribution of the size at division,  $P(s_d/\langle s_d \rangle|s_b)$ , for all the experiments. Figure S6 depicts  $P(\Delta_d|s_b)$ 



### Table of statistics

	panel A	panel B		
growth condition	width change from birth to division in µm	linear fit slope	% of width change over 20% of length change	width CV (%)
glycerol	-0.024	-0.019	0.4	9.03
sorbitol	-0.013	-0.079	1.6	5.18
glucose	-0.026	-0.037	0.7	5.03
glucose + 6. a.a.	-0.046	-0.031	0.6	6.05
glucose + 12 a.a.	-0.036	-0.06	1.2	4.22
synthetic rich	-0.015	-0.034	0.7	5.84
TSB	-0.006	-0.054	1.1	6.11

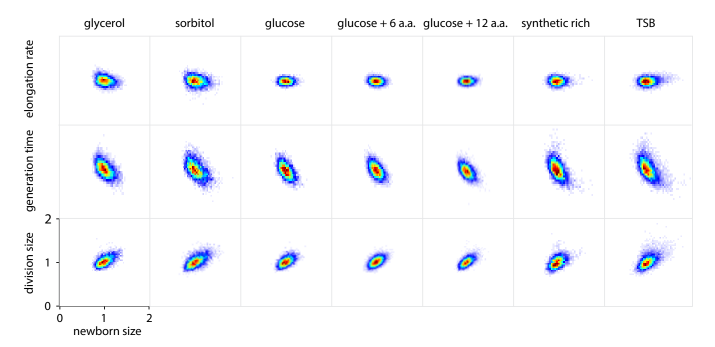
**Figure S4.** The behavior of the width of *E. coli* cells in the steady states at given growth conditions. (A) The width remains constant throughout the cell cycle, from birth to division. (B) The absence of correlation between the length at birth (rescaled by its mean value) of the cells and their width (rescaled by its mean value). The resulting correlation between width and length of the cells is negligible and smaller than the coefficient of variation of the width itself.

Table SII A numerical comparison of the variation of the cell width with respect to the newborn size and to the age of the cells. The second column (data from panel A) shows the change of the average width from birth to division, based on a linear fit to the data. The columns three and four (data from panel B) show the dependence of the cell width on the newborn length and the change of the average width normalized by the coefficient of variation (CV) of the newborn length, which is about 20%. The last column shows the coefficient of variation (CV) of the cell width in different conditions.

(top panels) and  $P(s_d|s_b)$  (bottom panels) in different growth conditions. We observe a consistent trend in all of the experiments: the distributions of  $\Delta_d$  are independent of the newborn cell size while the distributions of  $s_d$  clearly depend on  $s_b$ . The conditional distribution of the size at division,  $P(s_d/\langle s_d \rangle|s_b)$ , manifestly shifts as the newborn size  $s_b$  increases.

### D. Constant $\Delta$ evidence for E. coli K-12MG1655 and B/r strains

Using data from published work by Wang et al. [3], we have looked into the constancy of  $\Delta$  with respect to newborn size of the cells. The experiments have been performed in similar microfluidics device in LB media at 37°C. Figure 7 shows that the added size is indeed constant with respect to newborn size and that the distribution of  $\Delta/\langle\Delta\rangle$  is independent of newborn size.



**Figure S5.** Correlations between *E. coli* rescaled parameters in different growth conditions. The three rows show the rescaled (by their mean) elongation rate, generation time and size of the cell at division *vs* the size of the cell at birth. Note that the trends are similar for different conditions, strongly suggesting that the size control mechanism is independent of the growth conditions. The plot in Figure 1D of the main text is a combination of the data presented here.

### E. Correlation between growth rate $\lambda$ and added size $\Delta$

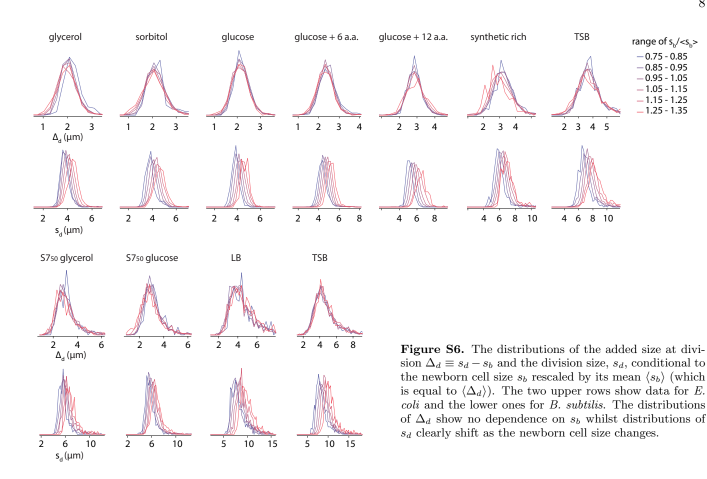
To justify use of the growth rate distribution  $\rho(\lambda)$  and the added size distribution  $\rho(\Delta)$  as two independent basic controls of physiology and size homeostasis, we measured their correlations. We confirm the two show negligible correlations (Fig. S8).

### F. Relaxation of atypical cell sizes to typical values

The  $\Delta$  model implies that if a group of cells is prepared with a size initially deviating from the population average value, their average size will approach the population average as generations elapse (see Section IV for details). To test this, we collected a subpopulation of the cells in each experiment, namely we sorted one group of large cells and one group of small cells. We then followed these subpopulations for 10 generations and monitored their average newborn size after each division. After 10 generations, the cell size of each group essentially converged to the average size ( $\langle s_b \rangle = \langle \Delta_d \rangle$ ) for the corresponding growth condition. In Figure 3b of the main text, we presented the convergence of the rescaled cell sizes for a combination of all experiments and we observed consistency with the  $\Delta$  model. Figure S9 shows the convergence of the size for each growth condition separately. Blue and red markers/lines indicate the large and small subpopulations, respectively. Markers with error bars are the results of the experiments; solid lines show the theoretical prediction of the  $\Delta$  model, which will be presented in Section IV.

### G. Distributions of the various growth parameters

Growth parameters, such as elongation rate, generation time, and size at birth and division, have some variability over the populations in a given growth condition. Figure 4 in the main text showed the remarkable collapse of the various distributions once the variables are rescaled by their mean. Here we present in Figure S10 the probability distributions of the same parameters yet with no rescaling by their mean. The table in Figure S10 reports the numerical values of the mean, the standard deviation, and the coefficient of variation (CV). The standard deviation for all the parameters increases as the mean increases. Conversely, the CV does not show any particular dependence



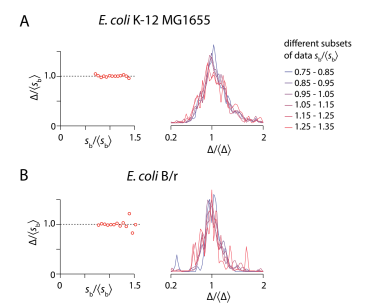
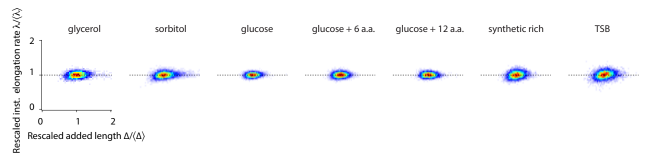
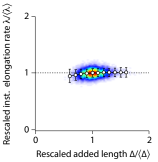


Figure S7. Data from E. coli strains K-12 MG1655 and B/r growth in steady-state in LB media at 37°C shows that the added size is indeed constant with respect to newborn size and that the distribution of  $\Delta/\langle \Delta \rangle$  is independent of newborn size

on the mean. As discussed in more detail in Section V, this behavior is consistent with a scale-invariant form for the probability distributions. An additional prediction is that higher-order moments of order p normalized by the power pof the mean should be a constant (dependent on p) with respect to the growth conditions. Such behavior is consistent with the data, as shown in Figure S11.



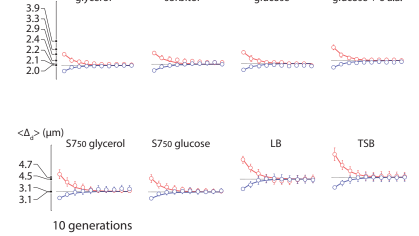


glycerol

 $<\Delta_d> (\mu m)$ 

Figure S8. Independence between the growth rate and the added size  $\Delta$ . Top panels show the scatter plots for individual experiments and the bottom left panel shows the average of the experiments binned data. The white marker and the error bars show mean and standard deviation in each bin. This justifies using  $\rho(\lambda)$  and  $\rho(\Delta)$  as two independent input parameters for self-consistency check of the  $\Delta$  model for cell-size control and homeostasis.

glucose + 12 a.a.



glucose

glucose + 6 a.a.

sorbitol

Figure S9. The  $\Delta$  model ensures that fluctuations in the cell size are controlled. We followed a subpopulation of big (red lines) and small (blue lines) cells for 10 generations. In each growth condition, the average size of each subpopulation converges to the average value for the whole population in the corresponding growth condition. The top row shows data for  $E.\ coli$  and the bottom row shows  $B.\ subtilis$  data. Markers with error bars show experimental data while solid lines show the theoretical predictions for the  $\Delta$  model discussed in Section IV.

synthetic rich

TSB

### H. Autocorrelations

Autocorrelation functions can reveal repeating patterns or memory in a series of data and the decay of memory across generations. The autocorrelation function of various parameters of interest, including elongation rate, generation time, newborn size, division size, and added size, are presented in Figure S12. The corresponding theoretical predictions for the  $\Delta$  model are discussed in the Section IV.A.

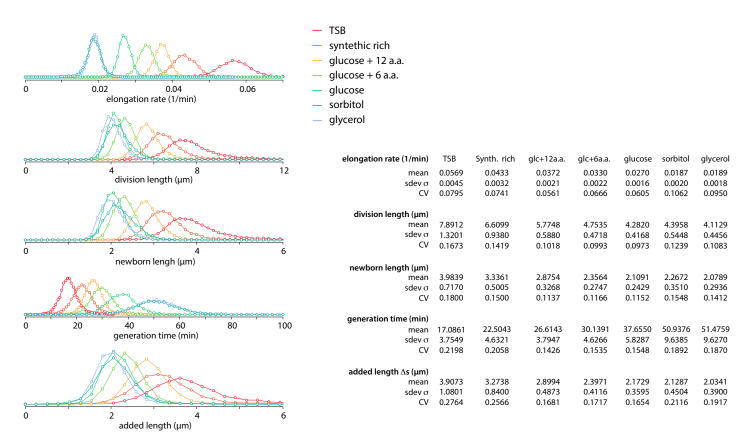


Figure S10. Normalized distributions of various *E. coli* parameters in different growth conditions. The table on the right shows the mean, the standard deviation and the coefficient of variation (CV) of the quantities plotted on the left.

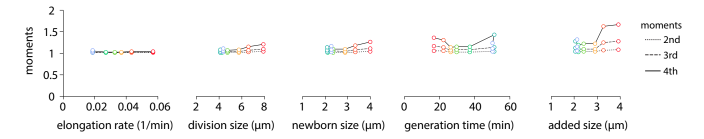
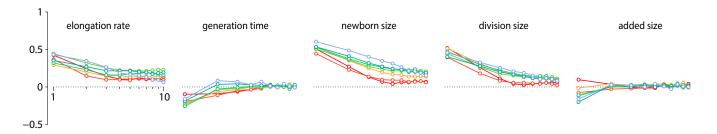


Figure S11. Higher-order moments of the distributions of various E. coli parameters. The moments of order p (p = 2, 3, 4), divided by the power p of the mean, are plotted as a function of the mean. For a scale-invariant distribution, those normalized moments should be constant with respect to the value of the mean, which is consistent with the data. Deviations observed for the fourth-order moment are due to partial statistical convergence, as witnessed by their increase moving from the left to the right graphs (the direction of increasing fluctuations, see Figure S10.)

### III. THEORY: A SUMMARY OF PREVIOUS MODELS FOR THE CELL SIZE CONTROL

Mechanisms previously proposed in the literature for the control of the cell size can be grouped into the following categories: (A) the control depends directly on the size of the cell ("sizer"); (B) the control is based on the age of the cell ("timer"); (c) combinations of the two previous controls. We briefly review the various mechanisms below. We also compare their predictions with the experimental data presented in the main text, illustrating the limitations of the existing models and the necessity of the  $\Delta$  model discussed later, based on the control of the added size.



**Figure S12.** The autocorrelations of various *E. coli* parameters across the generations. The first point (with the unit value on the abscissae) refers to the correlation among mother and daughters; correlations are measured up to 10 generations, when they have essentially vanished.

### A. The "sizer"

The main assumption of the model is that cells divide after reaching a threshold size, which is drawn from a probability distribution that depends on growth conditions. This mechanism explains by its very definition the observation that cells have a well-defined distribution of sizes. However, as shown below, its specific predictions are not supported quantitatively by our experimental data.

As shown by Figure S4, E. coli cells keep their width roughly constant during their growth and elongate along the axis of the rod. We shall denote the size of the cells along the axis by s and assume  $\dot{s}=g(s)$ , where the dot indicates the time derivative. Sizes of the cells at birth and division are denoted by  $s_b$  and  $s_d$ , respectively. The number of individuals in a population of bacteria having size s at time t is denoted n(s,t). The sizer mechanism posits that the rate of division of the cells depends on their size s only. The equation that governs the evolution in time of n(s,t) reads:

$$\partial_t n(s,t) + \partial_s (g(s) \, n(s,t)) = -g(s) \, \gamma(s) n(s,t) + 4g(2s) \gamma(2s) n(2s,t) \,. \tag{1}$$

The left-hand side in (1) is the total time-derivative, with the s-derivative drift term accounting for the elongation of the cells. The right-hand side of (1) stems from the division of cells and  $\gamma(s)$  is the so-called splitting rate function [see, e.g., [6]]. The function is defined as the local Poisson rate of cell division, i.e.,  $\gamma(s) ds$  is the probability that a cell of size s divides while growing from s to s+ds. Therefore, the probability that a cell with initial size  $s_b$  has not divided while reaching the size  $s_d$  is  $\exp\left[-\int_{s_b}^{s_d} \gamma(s') ds'\right]$ . Finally, the conversion of the division rate from unit size to unit time involves the factor |ds/dt| = g(s), which justifies its presence in the first term (of loss) on the right hand side of (1).

The second term in the right-hand side of (1) describes the gain in the number of cells in the size range (s, s + ds). The gain originates from those bacteria in the size range (2s, 2s + 2ds) which divide and thus halve their size (fluctuations in the size of the offsprings are neglected here for simplicity but will be discussed later). The structure of the second term is thus analogous to the previous one, with s replaced by s in the arguments of the functions. The factor 4 arises from the product of two factors 2: the first is due to the s range of dividing cells and the second is due to division producing two offsprings.

Equation (1) has a long history, which goes back at least to Ref. [7], where the basic formalism was introduced. The formalism was further developed and expanded in [8] and then utilized in a series of papers and books that include [6; 9; 10; 11; 12; 13]. Additional features, such as e.g. fluctuations in the rate of elongation or in the sizes of the offsprings due to the positioning of the septum can be included in the formalism [see, e.g., the Supplementary Material in [13]], but will not be needed here.

The solution to the equation (1) depends on the specific form of the splitting rate  $\gamma(s)$  and the elongation law g(s). It is possible though to extract a few general relations that we proceed to derive.

First, integrating (1) over s, we obtain the equality

$$\partial_t \int ds \, n(s,t) = \int ds \, g(s) \, \gamma(s) n(s,t) \,, \tag{2}$$

where the right-hand side is the total number of individuals which divide in the time interval [t, t + dt]. For the rate

of growth of the number of individuals in the population, we obtain then

$$\partial_t \ln \int ds \, n(s,t) = \int ds \, g(s) \, \gamma(s) \frac{n(s,t)}{\int ds \, n(s,t)} \equiv \langle g(s) \gamma(s) \rangle \,. \tag{3}$$

At long times, we expect from the very definition of the model that  $n(s,t)/\int ds \, n(s,t)$  will reach a stationary distribution that we denote by  $\rho_n(s)$ . The corresponding rate of growth of the population is denoted  $k = \int ds g(s) \gamma(s) \rho_n(s)$ . Second, multiplying (1) by s and integrating, we can obtain for the average size

$$\partial_t \langle s \rangle = \langle g(s) \rangle - \langle g(s) \gamma(s) \rangle \langle s \rangle.$$
 (4)

In the stationary state, the two terms on the right-hand side of (4) will balance. Namely, for an exponential elongation  $g(s) = \alpha s \ln 2$ , it follows from (4) that the consistency condition  $\langle s \gamma(s) \rangle = 1$  is satisfied. The factor  $\ln 2$  in g(s) is introduced to conform to the microbiology convention of cells elongating as  $s(t) = s_b 2^{\alpha t}$ .

Finally, multiplying (1) by  $s^q$  and integrating, we obtain a series of relations for higher-order moments. In the stationary state and for an exponential elongation  $g(s) = \alpha s \ln 2$ , the relations read  $(q-1)\langle s^q \rangle = (1-2^{1-q})\langle s^{q+1}\gamma(s) \rangle$ , where the integer q > 1 and we have made use of (4).

### 1. Comparison and consistency with experiments

The analytical solution of (1) is not available for generic forms of the elongation rate g(s) and of the splitting rate function  $\gamma(s)$ . Imposing a sharp maximum value S for the size distribution, i.e.  $n(s > S, t) \equiv 0$ , permits to patch the solution and have it in closed form (see the SI in [11] for details) yet the resulting expressions are not particularly transparent. Therefore, we decided to take a more pragmatic approach: g(s) and  $\gamma(s)$  are extracted from experimental data, e.g. from the distribution of the sizes at division and of elongation rates, and we use them to simulate the cell size control process at the level of individual cells. We can then compare additional statistical observables, other than those used to calibrate the model, to assess the consistency and validity of the sizer mechanism for the size control.

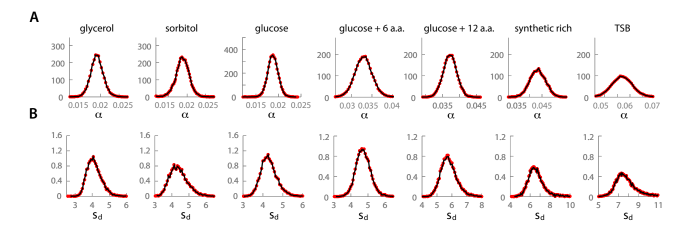


Figure S13. Calibration of the "sizer" model for the control of the cell size. Each cell's elongation rate is a random variable independently drawn from the experimental  $E.\ coli$  distribution,  $\rho_{\alpha}^{ex}(\alpha)$ , shown in red in panel (A) for the seven different growth conditions discussed in the Section I.A (in the order of increasing growth rate: glycerol, sorbitol, glucose, glucose + 6 a.a., glucose + 12 a.a., synthetic rich, TSB). Correlations among the elongation rates of mothers and siblings are weak and not taken into account; we verified that results are not modified by including them. The splitting rate  $\gamma(s)$  is computed from the distribution of the size of the cells at division  $\rho_d^{ex}(s_d)$  as detailed in the text (see eqs. (5) and (6)). The experimental distributions are shown in red in panel (B) for the same growth conditions as for the elongation rate. The curves in black are the results of numerical simulations of the model detailed in the text. Their agreement with the experimental curves confirms that the parameters (distribution of the elongation rate  $\alpha$  and the splitting rate function  $\gamma$ ) of the model are appropriately calibrated.

Specifically, the calibration of the model goes as follows. As for the elongation rates, experimental data are consistent with an exponential law,  $g(s) = \alpha s \ln 2$ , where the elongation rate  $\alpha$  can vary from cell to cell. The values

of  $\alpha$  that we use in the simulations are drawn directly from the experimental distribution,  $\rho_{\alpha}^{ex}(\alpha)$  shown in Figure S13 for *E. coli*. Draws are independent for different bacteria. Furthermore, experimental data indicate that the correlation between the elongation rates of mother and daughter cells is weak. We therefore assume that the elongation rate  $\alpha$  for each cycle of elongation of a bacterium is drawn independently of its past and future cycles of elongation.

As for the splitting rate function  $\gamma(s)$ , we compute it from the the experimental size distributions  $\rho_d^{ex}(s_d)$  and  $\rho_b^{ex}(s_b)$  of the sizes at division and birth, respectively. From the very definition of the splitting rate  $\gamma(s)$  we have at the steady state the following relation

$$\rho_d(s_d) = \gamma(s_d) \int_0^{s_d} \rho_b(s_b) e^{-\int_{s_b}^{s_d} \gamma(s') \, ds'} \, ds_b \,. \tag{5}$$

Indeed,  $\rho_b(s_b)$  is the probability that a newborn cell has size  $s_b$ , the exponential term is the probability that the cell will not divide up to  $s_d$  and, finally,  $\gamma(s_d) ds_d$  is the probability of division in the range  $(s_d, s_d + ds_d)$ . The integral runs over all possible values of  $s_b$  respecting the constraint  $s_d \geq s_b$ .

Using the experimental distributions for  $\rho_d$  and  $\rho_b$ , we can extract the splitting rate function  $\gamma$  from (5). In practice, the calculation is simplified by remarking that the distributions at division and birth are well separated and very weakly overlap in their tails, i.e.  $\gamma$  is small where  $\rho_b$  is appreciable. A good approximation to (5) is then to assume  $\rho_b(s_b) = \delta(s_b)$ , which gives

$$\rho_d(s_d) = -\frac{d}{ds_d} e^{-\int_0^{s_d} \gamma(s') \, ds'} \quad \Rightarrow \quad 1 - \int_0^{s_d} \rho_d(s') \, ds' = e^{-\int_0^{s_d} \gamma(s') \, ds'} \quad \Rightarrow \quad \gamma(s_d) = \frac{\rho_d(s_d)}{1 - \int_0^{s_d} \rho(s') \, ds'}, \tag{6}$$

where we first used that the cumulative distribution and its density are related by  $\rho_d(s_d) = -d \left[1 - \int_0^{s_d} \rho_d(s') \, ds'\right] / ds_d$  and then took a logarithm of the second equality and derived it with respect to  $s_d$ . The final approximation coincides with the expression for  $\gamma$  given in [6]. Experimental data for the size at division are well reproduced by the approximation (6), as shown in Figure S13.

We now proceed to simulate the dynamics of a growing population using the distribution of the elongation rates  $\alpha$  and the splitting function  $\gamma(s)$  extracted from the experimental data as we just described. Each cycle of elongation of a cell proceeds at the constant (random) rate  $\alpha$  and division occurs with the Poissonian rate  $\gamma(s)$ . After an initial transient time period, distributions for the various observables reach a stationary form, as expected. The resulting distributions for the final size  $s_d$  and  $\alpha$ , which we used to calibrate the model, are shown in Figure S13 for the case of  $E.\ coli.$  Similar curves are obtained for  $B.\ subtilis$  (not shown). Non-trivial tests for the sizer mechanism of control are provided by comparing the results of numerical simulations and experiments for additional observables.

The distribution for newborn sizes  $s_b$  is shown in Figure S14. The agreement is good, yet not independent of the calibration process itself since the two distributions of the size at birth and at division are simply related to each other as shown in panel (B) of the figure. A germane test of the model is provided by the distributions of the added size  $\Delta_d = s_d - s_b$  and of the generation time  $\tau_d = \log_2(s_d/s_b)/\alpha$  for each cell during the growth of the colony. Results obtained by simulations are compared with E. coli experimental data in Figure S15: the former are found to have fatter tails than the latter. The disagreement can be intuitively traced back to the fact that the sizer mechanism posits that division is essentially controlled by the cell size only. Indeed, given an initial size  $s_b$  the probability of division in the interval  $(s_d, s_d + ds_d)$  is  $\propto -d \exp\left[-\int_{s_b}^{s_d} \gamma(s') \, ds'\right]/ds_d$  [see (5)] and the dependency on  $s_b$  is weak, as we discussed for the derivation of (6). Conversely, experimental data show that the dependency between the initial and the final size of a cell is strong, as clearly illustrated by the shifts with  $s_b$  of the conditional probabilities  $P(s_d|s_b)$  shown by experimental data presented in the Figure S6. Discrepancies in Figure S15 are then intuited because the shift of  $P(s_d|s_b)$  with  $s_b$  reduces the contribution to the left (right) tails of  $\Delta_d = s_d - s_b$  and  $\tau_d$  from those cells whose initial size is larger (smaller) than the average.

A final check that the dependency of the division rate on  $s_b$  is indeed the crucial element missing in the sizer mechanism is provided in Figure S16. Picking randomly  $s_b$  and  $s_d$  from the two distributions  $\rho_b^{ex}(s_b)$  and  $\rho_d^{ex}(s_d)$  (with the only constraint  $s_d \geq s_b$  and no additional correlations) we show that the sizer model predictions are recovered. In conclusion, the sizer model ensures a stationary distribution of the cell size, yet its quantitative predictions disagree with the experiments. Discrepancies are due to the fact that the sizer mechanism neglects the role of the size at birth  $s_b$  on the control of the cell size at division  $s_d$ .

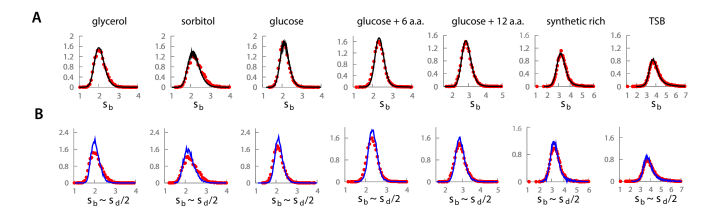


Figure S14. The "sizer" model for the control of the cell size reproduces some features of the experimental data but not all (see Figures S15 and S16). In panel (A), the model calibrated as in Figure S13 is simulated numerically and the distributions of the newborn cell size  $s_b$  are reported and compared to E. coli experimental data for the same growth conditions as in Figure S13. Black curves refer to numerics while red dots are the experimental data; the agreement is excellent. However, the agreement is strongly related to the calibration process since  $s_b \simeq s_d/2$ . Additional noise due to the fluctuations in the halving of the size of the daughter cells and the positioning of the septum is relatively weak, as shown in panel (B). The panel shows the experimental distributions of  $s_b$  (red dots) on top of the experimental distribution of the quantity  $s_d/2$  (blue lines) showing their strong overlap.

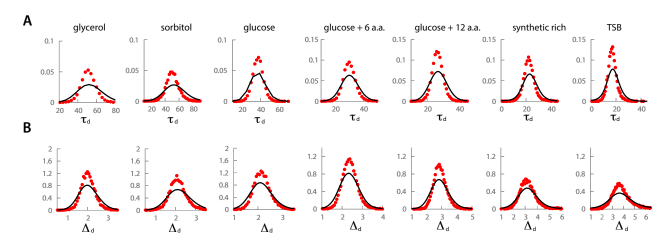


Figure S15. Test of the "sizer" model for the control of the cell size. The model calibrated as in Figure S13 is simulated numerically and we report the distribution of the generation time  $\tau_d = \log_2(s_d/s_b)/\alpha$  and of the added size  $\Delta_d = s_d - s_b$  for the case of E. coli. Black curves refer to numerics and red dots to experiments, respectively. The seven panels in (A) show the curves for the generation time  $\tau_d$  for all the growing conditions while panel (B) presents the distribution of the  $\Delta_d$ . Tails in the experimental data are significantly less pronounced for the reasons discussed in the body of the text. A similar disagreement is observed for B. subtilis.

### B. The "timer"

The "timer" mechanism posits that cell division is controlled by the age of the cell, i.e. the time elapsed since its birth. The state of cells is therefore described by their size s and their age  $\xi$  and the corresponding number of cells at time t is denoted by  $n(s, \xi, t)$ . The equation for the evolution of n reads:

$$\partial_t n(s,\xi,t) + \partial_s \left[ g(s)n(s,\xi,t) \right] + \partial_\xi n(s,\xi,t) = -\gamma(\xi)n(s,\xi,t); \quad n(s,0,t) = 4 \int \gamma(\xi')n(2s,\xi',t) \,d\xi'. \tag{7}$$

The left hand side is the total time derivative: the drift in s is due to the elongation of cells  $\dot{s} = g(s)$  while the drift in  $\xi$  is due to the aging of cells  $\dot{\xi} = 1$  (the discontinuity in  $\xi$  occurring at division will be addressed momentarily).

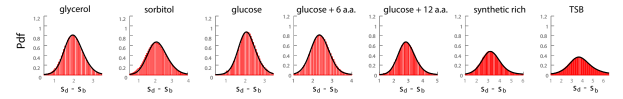


Figure S16. The "sizer" model is inconsistent with experimental data because it neglects the effects of the initial size  $s_b$  onto the control of the size  $s_d$  at division of the cells. The point is directly demonstrated by choosing randomly  $s_b$  and  $s_d$  from their respective experimental distributions  $\rho_b^{ex}(s_b)$  and  $\rho_d^{ex}(s_d)$  for E. coli. The only constraint we impose in the random draw is  $s_d \geq s_b$  and no additional correlations are retained. The resulting distributions for the "scrambled" added size  $s_d - s_b$  (shown in red) are indeed in agreement with the sizer model results (black curves).

The right hand side of the first equation in (7) is the loss term due to the division of cells. The Poisson splitting rate function  $\gamma$  depends now on the age of the cell,  $\xi$ . Furthermore, since  $\dot{\xi} = 1$  there is no additional factor coming from the conversion of the rate of division from unit age to unit time (see (1)). From the definition of the splitting rate it follows that

$$\rho_{\tau_d}(\tau) = \gamma(\tau) e^{-\int_0^{\tau} \gamma(\xi') d\xi'} \Rightarrow \gamma(\tau) = \frac{\rho_{\tau_d}(\tau)}{1 - \int_0^{\tau} d\xi' \rho_{\tau_d}(\xi')},$$
 (8)

where  $\rho_{\tau_d}(\tau)$  is the probability density for the generation time  $\tau_d$  of a given cell and the derivation proceeds as for (6). Finally, the last equation in (7) is the boundary condition that accounts for newborn cells having all the same age  $\xi = 0$ , irrespective of their size 2s that gets halved. The integral  $\int d\xi \, \gamma(\xi) n(2s, \xi, t)$  represents the total number of cells that divide in the unit time; as in (1), the factor 4 is the product of the factor 2 resulting from the 2 ds width of the range of dividing cells and the factor 2 due to division producing two newborn daughters cells.

The dynamics of the age of the cells is independent of their size, as it can be easily seen by integrating (7) over s. The resulting equations for the marginal distribution  $n(\xi,t) = \int n(s,\xi,t) ds$  reads

$$\partial_t n(\xi, t) + \partial_{\xi} n(\xi, t) = -\gamma(\xi) n(\xi, t); \quad n(0, t) = 2 \int d\xi \, \gamma(\xi) \, n(\xi, t). \tag{9}$$

It is also immediate to integrate (7) over s and  $\xi$ , obtaining for the rate k of growth of the population

$$\partial_t \int n(s,\xi,t) \, ds \, d\xi = \int \gamma(\xi) n(s,\xi,t) \, ds \, d\xi = \int \gamma(\xi) n(\xi,t) d\xi \,, \tag{10}$$

which depends on the marginal distribution  $n(\xi,t)$  only. The well-known solution [14] to (9) is  $n(\xi,t) = e^{kt}\tilde{n}(\xi)$ , where

$$\tilde{n}(\xi) = Ae^{-\int_0^{\xi} \gamma(\xi') \, d\xi' - k\xi}; \quad 2k \int_0^{\infty} e^{-k\xi} e^{-\int_0^{\xi} \gamma(\xi') \, d\xi'} \, d\xi = 1.$$
(11)

The first equation is simply obtained by integrating the differential equation in (9) and the constant A depends on the initial size of the population. The second relation is obtained from the boundary condition in (9) by an integration by parts. A similar integration by parts also allows to check that the second equation in (11) is equivalent to the growth rate relation  $k \int \tilde{n}(\xi) d\xi = \int \tilde{n}(\xi) \gamma(\xi) d\xi$  derived in (10).

As for the dynamics of the size of cells, we can obtain the equation for the mean size multiplying (7) by s and integrating over s and  $\xi$ . The resulting expression reads

$$\partial_t \langle s \rangle = \langle g(s) \rangle - k \langle s \rangle, \tag{12}$$

where  $\langle s \rangle \equiv \int s \, n(s,\xi,t) \, ds \, d\xi / \int n(s,\xi,t) \, ds \, d\xi$  and similar definitions apply for other averages. For a stationary state, the left-hand side is required to vanish, i.e. elongation should balance the growth. In the case of linear elongation, g(s) = const. a stationary distribution is reached and  $\langle s \rangle = \text{const.} / \langle \gamma(\xi) \rangle$ , where we used  $k = \langle \gamma(\xi) \rangle$ . Conversely, for exponential elongation  $g(s) = \alpha s \ln 2$ , a stationary distribution obtains only for the special choice  $\alpha \ln 2 = k$ . If the equality is not satisfied, then the average size grows to infinity or decay to zero exponentially fast (see below).

The lack of control of the size of the cells by the timer mechanism was remarked in [15; 16] and it is intuitively understood by considering the sizes at birth  $s_b^{(n)}$  and  $s_b^{(n+1)}$  over two consecutive generations n and n+1. For an

exponential elongation,  $\log_2 s_b^{(n+1)} - \log_2 s_b^{(n)} \simeq \alpha \tau_d - 1$ , where  $\tau_d$  is again the generation time and  $\alpha$  is the elongation rate for the bacterium in the n-th generation. Exponential elongation of the size of the cells makes that the average value of  $\alpha \tau_d$  must be precisely tuned in order to avoid a systematic drift of  $\log_2 s_b$ . Notice that even in the absence of drift, the long term behavior of  $\log_2 s_b$  will be analogous to a random walk (assuming that the values of  $\alpha \tau_d$  fluctuate and decorrelate over the generations). Therefore, the variance of the size of the cells will grow across the generations and no effective control of the size of the cells is achieved by the timer mechanism. Diluting bacteria by washing them out, e.g. a term  $-Dn(s,\xi,t)$  is added to the equation (7), will not modify the previous conclusion unless dilution is coupled to the size, i.e. the dilution rate depends on s.

In practice, the elongation rate is deviating from a linear behavior at very small and very large sizes so that the logarithm of the size will not go to zero or diverge to infinity. However, its behavior will depend very sensitively on the details of the elongation law at very small and large sizes [13; 17] and the resulting size distributions are generally too wide (data not shown). Most importantly, the timer mechanism disagrees with our experimental data in that the conditional distribution  $P(\tau_d|s_b)$  of the generation time  $\tau_d$  vs the initial size of cells  $s_b$  should be independent of  $s_b$ . Conversely, data shown in the main text indicate a clear dependency on  $s_b$ .

### C. Mixed models

Diverse combinations of sizer and timer mechanisms are conceivable. A well-known instance is the bilinear model proposed in [18]. The proposed dynamics is summarized as follows: cells grow linearly in time g(s) = u until they reach a threshold size  $s_{\Lambda}$ . The threshold is argued to be universal with respect to growth conditions and its value  $s_{\Lambda} \simeq 2.78 \,\mu m$  to be twice the minimal possible length for  $E.\ coli$  cells. After reaching  $s_{\Lambda}$ , cells keep growing for a fixed amount of time ( $\sim 20$  mins) at a velocity 2u (see [19; 20; 21] for a discussion of bilinear vs exponential elongation). The velocity u depends on the growth conditions and the last 20 mins phase accounts for the dependency of the distribution of the final size  $s_d$  on u, and thus on the growth conditions. Ref. [18] sharply poses relevant questions and deserves all the influence it had in the field. However, the specific mechanism which was proposed is not supported by modern experimental data that allow for better statistics. In Figure S17, the size  $s_d$  and the generation time  $\tau_d$  at division vs the initial size  $s_b$  are presented and the disagreement between the predictions and the data is manifest.

More recently, other combinations of age and size control have been proposed. The intuitive motivation comes from plots like Figure 1D in the main text, where a behavior intermediate between a timer and a sizer is observed. The most recent proposition is [12], where a control mechanism operating concertedly (and not sequentially as in [18]) on the size and the age of the cell is discussed. Namely, no major inconsistency with experimental data (for a single growth condition) is observed by introducing a model where the splitting rate function  $\gamma(s,\xi)$  depends jointly on the size and on the age of the cells and by best fitting the function to the data.

### IV. THEORY: THE $\Delta$ MODEL OF CONTROL

The  $\Delta$  model that we propose and discuss in the main text posits that the mechanism of control involves a single variable,  $\Delta$ , the size added to the cells since their birth. The purpose of this Section is to provide further details on the model and its comparison to experimental data.

As in the previous Section, we shall denote by s the size of cells along the elongating axis of the rod. We shall assume ds/dt = g(s) and the width of the cell is roughly constant. We indicate by  $s_b$  and  $s_d$  the size of cells at birth and division. If s(t) is the size of a cell at the current time t, its added size is denoted  $\Delta(t) = s(t) - s_b$ . The density of cells  $n(s, \Delta, t)$  having size s and added size  $\Delta$  obeys the following continuity equation

$$\partial_t n(s, \Delta, t) + \partial_s \left[ g(s) n(s, \Delta, t) \right] + \partial_\Delta \left[ g(s) n(s, \Delta, t) \right] = -\gamma(\Delta) g(s) n(s, \Delta, t); \tag{13}$$

$$g(s)n(s,0,t) = 4g(2s) \int_0^\infty \gamma(x)n(2s,x,t) \, dx.$$
 (14)

The left-hand side in (13) is the total time-derivative and the two drift terms are due to the elongation of the cells, i.e. ds/dt = g(s) and  $d\Delta/dt = g(s)$ . As in the previous Sections on the sizer and the timer, the right hand side accounts for the division of the cells. The Poissonian splitting rate function  $\gamma(\Delta)$  depends now on the added size  $\Delta$ . Proceeding as for (6) and (8), we obtain the relation

$$\rho_{\Delta_d}(\Delta) = \gamma(\Delta) e^{-\int_0^\Delta \gamma(x) \, dx} \Rightarrow \gamma(\Delta) = \frac{\rho_{\Delta_d}(\Delta)}{1 - \int_0^\Delta dx \, \rho_{\Delta_d}(x)}, \tag{15}$$

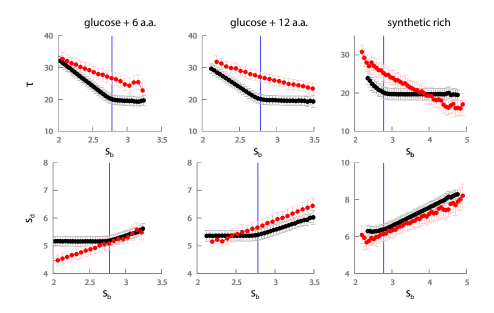


Figure S17. Comparison of bilinearity (black points) with E. coli experimental data (red points) for the fast growth conditions glucose + 6 a.a., glucose + 12 a.a. and synthetic rich. The behavior of the size  $s_d$  and the generation time  $\tau_d$  at division vs the initial size  $s_b$  are presented. A change of behavior should take place [18] at the size  $s_\Lambda$  (indicated by the vertical blue line) but data do not support any such transition. In particular, for  $s_b < s_\Lambda$  the division size  $s_d$  should be independent of  $s_b$  and for  $s_b > s_\Lambda$  the generation time  $\tau_d$  should be independent of  $s_b$ . Neither one of these predictions is observed.

between  $\gamma(\Delta)$  and the distribution  $\rho_{\Delta_d}(\Delta)$  for the size added at division ( $\Delta_d = s_d - s_b$ ) of individual cells. By individual cells we mean that cells should be weighted equally, tracking them individually and avoiding known bias effects related to the speed of reproduction [14]. As in the equation (1), the conversion of the rate of division to unit time involves the Jacobian  $|d\Delta/dt| = g(s)$ , that appears then in the right hand side of (13). Finally, (14) is the boundary condition that accounts for cells having all  $\Delta = 0$  at birth, irrespective of their size 2s that gets halved. The integral  $\int \gamma(x) n(2s, x, t) dx$  represents the total number of cells that divide in the unit time; as in (1), the factor 4 is the product of the factor 2 resulting from the 2 ds width of the range of dividing cells and the factor 2 due to division producing two newborn daughters cells.

A series of relations analogous to those obtained for the sizer model can be derived from (13) for a general splitting rate  $\gamma$ . Integrating (13) over s and  $\Delta$  and using the boundary condition (14), we obtain

$$\partial_t \ln \int n(s, \Delta, t) ds d\Delta = \int g(s) \gamma(\Delta) \frac{n(s, \Delta, t)}{\int n(s, \Delta, t) ds d\Delta} ds d\Delta \equiv \langle g \gamma \rangle.$$
 (16)

Multiplying (13) by s and integrating over s and  $\Delta$ , we derive for the average size

$$\partial_t \langle s \rangle = \langle g(s) \rangle - k \langle s \rangle, \tag{17}$$

where the rate of growth of the population  $k = \langle g(s)\gamma(\Delta)\rangle$  from (16). For a linear elongation, g(s) = u, (17) gives for the size at equilibrium  $\langle s \rangle = 1/\langle \gamma(\Delta) \rangle$ . For an exponential elongation,  $g(s) = \alpha s \ln 2$ , (17) yields  $\alpha \ln 2 = k$ , i.e. the value  $\langle s\gamma(\Delta) \rangle = 1$  for the correlation between the size and the splitting rate function.

Finally, we can obtain the following series of relations for higher order moments at the steady state:

$$(1-m)\langle s^m \Delta^q \rangle - q\langle s^{m+1} \Delta^{q-1} \rangle = 2^{1-m} \delta_{q,0} \langle s^{m+1} \gamma(\Delta) \rangle - \langle s^{m+1} \Delta^q \gamma(\Delta) \rangle, \qquad m \ge 0, q \ge 0, \tag{18}$$

where we specified relations for the case  $g(s) = \alpha s \ln 2$  and we used  $k = \alpha \ln 2$  and  $\langle s\gamma \rangle = 1$  derived previously.

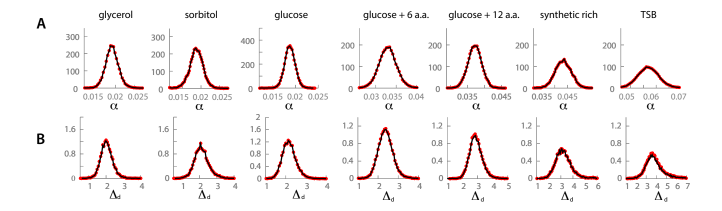


Figure S18. Calibration of the  $\Delta$  model for the control of the cell size. The elongation rate of the cells is an iid random variable drawn from the  $E.\ coli$  experimental distribution,  $\rho_{\alpha}^{ex}(\alpha)$  (red dots) in the plots of panel (A) for the seven different growth conditions presented in Figure S13. Correlations among elongation rates of mother and siblings are weak and thus not taken into account. We verified that results are not modified by including them. The black curves are the results of the numerical simulations. The splitting rate  $\gamma(\Delta)$  is computed as detailed in the text (see (15)) from the distribution  $\rho_{\Delta d}^{ex}(\Delta)$  of the increments at division  $\Delta_d = s_d - s_b$ . In panel (B) we show the experimental distributions (red dots) for the same growth conditions as in panel (A). The curves in black are the results of numerical simulations of the model detailed in the text. Their agreement with the experimental curves confirms that the parameters of the model are appropriately calibrated. Similar curves are obtained for  $B.\ subtilis$ .

### A. Comparison and consistency with experiments

The comparison with experimental data for the  $\Delta$  model proceeds as for the sizer mechanism. The elongation rate g(s) and the splitting function  $\gamma(\Delta)$  are extracted from experimental data, namely from the distribution of the sizes at division and the distribution of the elongation rates, and we use them to simulate the cell size control process at the level of individual cells. We then compare statistical observables alternative to those used for calibration, in order to assess the validity of the model.

The calibration of the model goes as follows. The elongation rate for individual bacteria is obtained by exponential fits of the experimental curves of size versus time. The probability densities  $\rho_{\alpha}^{ex}(\alpha)$  of the resulting elongation rates in the various growth conditions are shown in Figure S18. In our numerical simulations we draw values of  $\alpha$  randomly from  $\rho_{\alpha}^{ex}(\alpha)$ , neglecting (as for the sizer case) weak correlations between the elongation rates of mother and daughter cells. The splitting rate  $\gamma(\Delta)$  is computed using (15) with the probability of the size increments at division  $\Delta_d \equiv s_d - s_b$  read directly from the experimental data  $\rho_{\Delta d}^{ex}(\Delta)$  (see Figure S18).

The distribution of the elongation rates  $\alpha$  and the splitting function  $\gamma(\Delta)$  extracted from the experimental data as we just described are used to simulate the dynamics of a bacterial colony. Each cycle of elongation of a cell proceeds at the constant (random) rate  $\alpha$  and division occurs with the Poissonian rate  $\gamma(\Delta)$ , which depends on the size increment  $\Delta$  only. After an initial transient, distributions for the various observables reach a stationary form and the resulting numerical distributions for the added size at division  $\Delta_d = s_d - s_b$  and  $\alpha$  (used to calibrate the model) compare to the experimental ones as shown in Figure S18.

Distributions of size and age at division. A first test for the  $\Delta$  model is provided by the curves in Figure S19 showing the agreement of the distributions for the final size  $s_d$  and for the generation time  $\tau_d = \log_2(s_d/s_b)/\alpha$  (where the values of the various quantities are those of individual bacteria). The corresponding results for B. subtilis are shown in Fig. S20. Most importantly, the model by its very definition agrees with the  $s_b$ -independent curves of the conditional probability  $P(\Delta_d|s_b)$  shown in the main text.

Correlations of the size across generations. An additional test for the model comes from the correlations of the size among genealogically related cells. Some of the correlations below have been calculated in Ref. [22] by a slightly different procedure. Correlations between the added length of the mother  $\Delta_d^M$  and the daughter,  $\Delta_d^D$  are experimentally found to be small so we shall neglect them hereafter.

experimentally found to be small so we shall neglect them hereafter. Let us start by the correlation  $\mathcal{C}_{dd}^{(p)} \equiv \langle s_d^M s_d^{D(p)} \rangle$  between the size at division of the mother and its p-th generation descendant. For example, p=1 gives the correlation between the sizes at division of mother and daughter

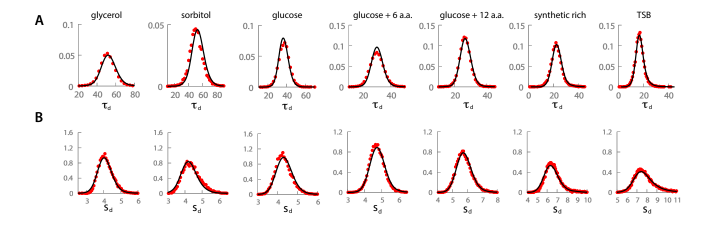


Figure S19. Test of the  $\Delta$  model for the control of the cell size. The model calibrated as in Figure S18 is simulated numerically and the *E. coli* distributions of the generation time  $\tau_d = \log_2(s_d/s_b)/\alpha$  and the size  $s_d$  at division of the cells are reported in panels (A) and (B), respectively, for the various growth conditions. Red dots refer to experiments while black curves are the numerics. The agreement of theoretical predictions with experimental data confirms the validity of the  $\Delta$  mechanism for the control of the cell size.

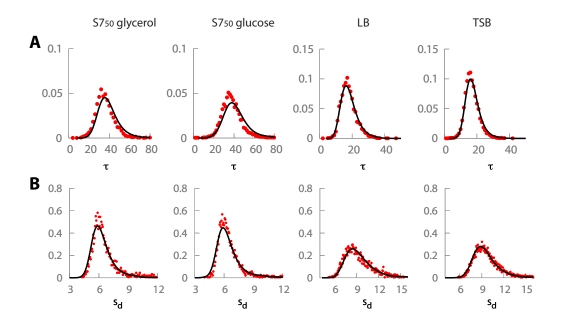


Figure S20. Test of the  $\Delta$  model for B. subtilis. As in Fig. S19, the distributions of the generation time  $\tau_d = \log_2(s_d/s_b)/\alpha$  and the size  $s_d$  at division of the cells are reported in panels (A) and (B), respectively, for the various growth conditions. Red dots refer to experiments while black curves are the numerical predictions.

cells. The decay of the correlation function is computed by the formula

$$\mathcal{C}_{dd}^{(p+1)} = \left\langle s_d^M s_d^{D(p+1)} \right\rangle = \left\langle s_d^M \left( \frac{s_d^{D(p)}}{2} + \Delta_d^{(p+1)} \right) \right\rangle = \frac{1}{2} \left[ \mathcal{C}_{dd}^{(p)} + \langle s_d \rangle^2 \right].$$
(19)

where we have used that  $\langle \Delta_d \rangle = \langle s_b \rangle = \langle s_d \rangle/2$ , as can be easily derived from  $s_d = s_b + \Delta_d$  and  $\langle s_b \rangle = \langle s_d \rangle/2$ . For the connected part of the correlation function  $C_{dd}^{(p)} \equiv C_{dd}^{(p)} - \langle s_d \rangle^2$ , we obtain then

$$C_{dd}^{(p)} \equiv \langle s_d^M s_d^{D(p)} \rangle - \langle s_d^M \rangle \langle s_d^{D(p)} \rangle = \sigma_{s_d}^2 / 2^p \quad \text{where} \quad \sigma_{s_d}^2 = \langle s_d^2 \rangle - \langle s_d \rangle^2 \,. \tag{20}$$

The comparison with the numerical simulations is shown in Figure S21. By similar arguments we can show that the correlation function for the size at birth  $C_{bb}^{(p)} = \sigma_{s_b}^2/2^p$  where the variance of the size at birth  $\sigma_{s_b}^2 = \langle s_b^2 \rangle - \langle s_b \rangle^2 \simeq \sigma_{s_d}^2/4$  because the size is (roughly) halved at division. The scaling is confirmed in Figure S21. Finally, the mixed correlation  $C_{bd}^{(p)} = C_{bb}^{(p)}$  because the added size is statistically independent of the initial size.

The positive correlation between the sizes across generations is quite intuitive to grasp. An ancestor cell bigger than the average will generate progeny that statistically relaxes to the average size, as we showed in the main text and the Figure S9. The fact that the added length is independent of the initial size makes that progeny will inherit only part of the ancestral size, which is successively halved as generations proceed. That accounts for the positive correlations and its 1/2 rate of decay across generations.

Notice that the previous results give for the Pearson correlation coefficient between the size at birth and at division:

$$\frac{\langle s_b \, s_d \rangle - \langle s_b \rangle \langle s_d \rangle}{\sigma_{s_b} \sigma_{s_d}} = \frac{C_{bd}^{(0)}}{\sigma_{s_b} \sigma_{s_d}} = \frac{\sigma_{s_b}}{\sigma_{s_d}} \simeq \frac{1}{2}, \tag{21}$$

which accounts for the behavior of experimental data observed in Figure 1D of the main text. It also follows from  $s_d = s_b + \Delta_d$  and  $s_b \simeq s_d/2$  that for the  $\Delta$  model

$$\sigma_{s_d}^2 = 4\sigma_{s_b}^2 = \frac{4}{3}\sigma_{\Delta_d}^2; \quad \langle s_d \rangle = 2\langle s_b \rangle = 2\langle \Delta_d \rangle. \tag{22}$$

We conclude that the coefficients of variation CV of the three quantities are

$$CV_{s_d} \equiv \frac{\sigma_{s_d}}{\langle s_d \rangle} \simeq CV_{s_b} \simeq \frac{1}{\sqrt{3}} CV_{\Delta_d} \,.$$
 (23)

Predictions are in excellent agreement with the experimental data presented in the main text. The coefficient of variation of  $s_b$  is actually slightly larger than  $CV_{s_d}$  before  $s_b$  is also affected by the noise in the positioning of the septum. We neglected here that source of noise because it is small; when included, it leads to  $CV_{s_b}$  being slightly larger than  $CV_{s_d}$ , as observed in the main text.

Correlations of generation times. Generation times  $\tau_d = 1/\alpha \log_2{(s_d/s_b)}$  involve a logarithm, which a priori is not obvious to treat. However, we can circumvent the problem using that the average of logarithms involve typical fluctuations around the mean and the observation that the coefficient of variation of  $s_b$  and  $\Delta_d$  are relatively small. We can then derive approximate expressions for those correlations. Neglecting for simplicity the small noise in  $\alpha$ , i.e.  $\alpha = \langle \alpha \rangle$  below, we expand  $\tau_d$  and  $1/\tau_d$  as

$$\tau_d = \frac{1}{\alpha \ln 2} \ln \left( 1 + \frac{\Delta_d}{s_b} \right) \simeq \frac{1}{\alpha} \left[ 1 + \frac{1}{2 \ln 2} \frac{\delta \Delta_d}{\langle \Delta_d \rangle} - \frac{1}{2 \ln 2} \frac{\delta s_b}{\langle s_b \rangle} \right]; \quad \frac{1}{\tau_d} \simeq \alpha \left[ 1 - \frac{1}{2 \ln 2} \frac{\delta \Delta_d}{\langle \Delta_d \rangle} + \frac{1}{2 \ln 2} \frac{\delta s_b}{\langle s_b \rangle} \right], \quad (24)$$

where  $\delta \Delta_d$  and  $\delta s_b$  denote fluctuations with respect to their respective mean values and we used  $\langle s_b \rangle = \langle \Delta_d \rangle$  (see (22)). Second-order terms will not be needed as they cancel out from the correlations we shall compute below.

From (24) we obtain for the mean, the variance and the coefficient of variation of  $\tau_d$ :

$$\langle \tau_d \rangle \simeq \frac{1}{\alpha}; \quad \langle \tau_d^2 \rangle - \langle \tau_d \rangle^2 \simeq \frac{1}{4\alpha^2 \ln^2 2} \left[ \frac{\sigma_{\Delta_d}^2}{\langle \Delta_d \rangle^2} + \frac{\sigma_{s_b}^2}{\langle s_b \rangle^2} \right] = \frac{1}{\alpha^2 \ln^2 2} \frac{\sigma_{s_b}^2}{\langle s_b \rangle^2} \quad \Rightarrow \quad CV_{\tau_d} = \frac{CV_{s_b}}{\ln 2} = \frac{CV_{\Delta_d}}{\sqrt{3} \ln 2}, \tag{25}$$

where we used (23) in the second equation. The ratios of the coefficients of variation are in excellent agreement with the experimental data presented in Fig. 4B of the main text.

Similarly, we can compute the mean value and the variance for  $1/\tau_d$ , which read:

$$\left\langle \frac{1}{\tau_d} \right\rangle \simeq \alpha \,; \quad \left\langle \frac{1}{\tau_d^2} \right\rangle - \left\langle \frac{1}{\tau_d} \right\rangle^2 \simeq \frac{\alpha^2}{\ln^2 2} \frac{\sigma_{s_b}^2}{\langle s_b \rangle^2} \,,$$
 (26)

and derive then

$$\frac{\left\langle \frac{s_b}{\tau_d} \right\rangle - \left\langle s_b \right\rangle \left\langle \frac{1}{\tau_d} \right\rangle}{\sigma_{1/\tau_d}^2} \simeq \frac{\left\langle s_b \right\rangle \ln 2}{2\alpha} \,. \tag{27}$$

The expression (27) was used in Figure 1C of the main text to fit the dependency of the initial size versus the number of divisions per hour, in given growth conditions. The fitting curve was written as  $s_b = c_1 \frac{1}{\tau_d} + c_2$  and it is verified that the constant  $c_1$  is then equal to the correlation (27).

We can also derive the correlation between  $s_b$  and the generation time  $\tau_d$  as

$$\langle s_b \tau_d \rangle - \langle s_b \rangle \langle \tau_d \rangle \simeq -\frac{1}{2\alpha \ln 2} \frac{\sigma_{s_b}^2}{\langle s_b \rangle} \quad \Rightarrow \quad \frac{\langle s_b \tau_d \rangle - \langle s_b \rangle \langle \tau_d \rangle}{\sigma_{s_b} \sigma_{\tau_d}} \simeq -\frac{1}{2} \,,$$
 (28)

where we used (25). The reason for the negative sign is quite intuitive. Indeed, the elongation at a size s proceeds at the rate  $\alpha s \ln 2$ , i.e. the longer the cell, the faster it grows. Therefore, if  $s_b$  is larger/smaller than the mean it will take less/more time to complete the addition of the size  $\Delta_d$  (independent of  $s_b$ ).

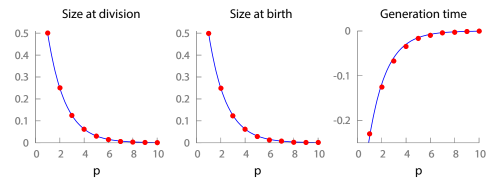


Figure S21. Correlations in the  $\Delta$  model. We simulate the process under the  $\Delta$  model and assume no correlation between the  $\Delta_d$  of the mother and its siblings. The connected correlation function  $\langle s_d^M s_d^{D(p)} \rangle - \langle s_d^M \rangle \langle s_d^{D(p)} \rangle$ , divided by its value  $\sigma_{s_d}^2$  for p=0, is plotted as a function of the generation p in panel (A). The line is the prediction derived in the text  $2^{-p}$  while the dots are numerical values. The corresponding correlation for the newborn size  $s_b$  is shown in panel (B). Finally, the connected correlation of the generation time defined in (29) as a function of the generations p is shown in panel (c). The best fit for the decay is the exponential behavior  $-0.43 \times 2^{-p}$ , confirming the behavior derived in the text.

Finally, we can explicitly compute the decay of the correlations among the division times across generations. We indicate by  $\tau_d^M$  the generation time of the mother and by  $\tau_d^{D(p)}$  the generation time of a p-th generation descendent. For instance, daughters correspond to p=1. We are interested in the behavior of the correlation

$$C_{\tau_d \tau_d}^{(p)} \equiv \frac{\left\langle \tau_d^M \tau^{D(p)} \right\rangle - \left\langle \tau_d^M \right\rangle \left\langle \tau_d^{D(p)} \right\rangle}{\sigma_{\tau_d}^2} \,, \tag{29}$$

Using (24), we can approximate the correlation by

$$C_{\tau_d \tau_d}^{(p)} \simeq -\frac{1}{\left[CV_{\Delta_d}^2 + CV_{s_b}^2\right]} \left\langle \left(\frac{\delta \Delta_d^M}{\langle \Delta_d^M \rangle} - \frac{\delta s_b^M}{\langle s_b^M \rangle}\right) \frac{\delta s_b^{D(p)}}{\langle s_b^{D(p)} \rangle} \right\rangle, \tag{30}$$

where we used that  $\delta \Delta_d^{D(p)}$  is independent of all other fluctuations in the  $\Delta$  model and the expression (25) of the variance  $\sigma_{\tau_d}^2$  in terms of the coefficients of variation. Note that the mean  $\langle s_b^{D(p)} \rangle$  is the usual value of the mean (unaffected by the mother's fluctuations) as corrections would yield higher-order corrections.

The size at birth of a p-th generation descendant is  $s_b^{D(p)} = \frac{\left(s_b^M + \Delta_d^M\right)}{2^p} + \sum_{k=1}^{p-1} \frac{\Delta_d^{D(k)}}{2^{p-k}}$ . Since the increments  $\Delta_d^{D(k)}$  are independent of the mother's fluctuations in the  $\Delta$  model, we conclude that

$$C_{\tau_d \tau_d}^{(p)} \simeq -\frac{CV_{\Delta_d}^2 - CV_{s_b}^2}{CV_{\Delta_d}^2 + CV_{s_b}^2} \times \frac{1}{2^p}$$
 (31)

Using (23), we obtain for the correlation between mother and daughters  $C_{\tau d\tau d}^{(1)} \sim -\frac{1}{4}$ . The behavior (31) is consistent with the experimental data in the Figure S12 and in the main text and with the results of numerical simulations in Figure S21 (small corrections to the constant prefactor are due to the noise in the positioning of the septum and fluctuations in the elongation rate  $\alpha$ ; we explicitly verified that in their absence the constant agrees indeed with (31)).

Anticorrelations between the generation times of the mother and its descendants are intuitive. Let us consider a given initial size for the mother. As its generation time becomes longer, the mother will divide bigger and will then tend to have descendants with a bigger newborn size. The size at birth and the generation time of a cell tend to be anticorrelated. This follows intuitively from the definition  $2^{\alpha\tau_d} = s_d/s_b = 1 + \Delta_d/s_b$  and the fact that the added size is independent of the initial size, i.e. it takes less to add the fixed amount  $\Delta_d$  if the cell elongates faster (see (28) and (32) for a more formal proof). Combining the two statements above we conclude that the division times of the mother and its descendants are anticorrelated.

Correlations involving exponentials of the generation time. We conclude this Section by computing some correlations involving exponentials of the generation time. The reason is that since  $2^{\alpha\tau_d} = s_d/s_b$ , correlations do not involve any logarithm and we can demonstrate anticorrelations without any hypothesis on the strength of the fluctuations. We first show that the initial size and  $2^{\alpha\tau_d}$  are anticorrelated in the  $\Delta$  model:

$$\langle s_b 2^{\alpha \tau_d} \rangle - \langle s_b \rangle \langle 2^{\alpha \tau_d} \rangle = \langle s_d \rangle - \langle s_b \rangle \left( 1 + \langle \Delta_d \rangle \left\langle \frac{1}{s_b} \right\rangle \right) = \langle s_b \rangle \left( 1 - \langle s_b \rangle \left\langle \frac{1}{s_b} \right\rangle \right) \le 0. \tag{32}$$

Here, we used  $\langle \Delta_d \rangle = \langle s_b \rangle = \langle s_d \rangle/2$ , and the inequality  $\langle x \rangle \langle 1/x \rangle \geq 1$  holding for any positive-definite random variable x (as it can be proved for example by the Cauchy-Schwartz inequality  $1 = \langle \sqrt{x} \sqrt{1/x} \rangle \leq \langle x \rangle^{1/2} \langle 1/x \rangle^{1/2}$ ).

The exponentials of the division times of the mother and its daughters are also anticorrelated. Indeed:

$$\left\langle 2^{\alpha^{M} \tau_{d}^{M}} 2^{\alpha^{D} \tau^{D}} \right\rangle = \left\langle \frac{s_{d}^{M}}{s_{b}^{M}} \frac{s_{d}^{D}}{s_{b}^{D}} \right\rangle = 2 \left\langle \frac{s_{d}^{D}}{s_{b}^{M}} \right\rangle = 2 \left\langle \frac{s_{d}^{M}/2 + \Delta_{d}^{D}}{s_{b}^{M}} \right\rangle = 1 + 3 \left\langle \Delta_{d} \right\rangle \left\langle \frac{1}{s_{b}} \right\rangle, \tag{33}$$

where we have used again that in the  $\Delta$  model, the added size  $\Delta_d$  is statistically independent of  $s_b$ . Subtracting then the disconnected contribution

$$\left\langle 2^{\alpha^{M}\tau_{d}^{M}}\right\rangle \left\langle 2^{\alpha^{D}\tau_{d}^{D}}\right\rangle = \left\langle \frac{s_{d}}{s_{b}}\right\rangle^{2} = \left(1 + \left\langle \Delta_{d}\right\rangle \left\langle \frac{1}{s_{b}}\right\rangle\right)^{2},\tag{34}$$

and using again  $\langle \Delta_d \rangle = \langle s_b \rangle$ , we finally obtain

$$\left\langle 2^{\alpha^{M}\tau_{d}^{M}}2^{\alpha^{D}\tau_{d}^{D}}\right\rangle - \left\langle 2^{\alpha^{M}\tau_{d}^{M}}\right\rangle \left\langle 2^{\alpha^{D}\tau_{d}^{D}}\right\rangle = \left\langle s_{b}\right\rangle \left\langle \frac{1}{s_{b}}\right\rangle \left(1 - \left\langle s_{b}\right\rangle \left\langle \frac{1}{s_{b}}\right\rangle\right) \leq 0. \tag{35}$$

### V. COLLAPSE OF THE PROBABILITY DISTRIBUTIONS AND SCALING FORMS

It was recently observed in [11] that intra- and inter-species distributions of body sizes have a universal form across many species. The universal form seems to be uniquely determined by the mean of the distribution, i.e., when the various distributions are rescaled by their mean, they tend to collapse onto a unique curve. This recent remark generalizes analogous, classical observations made for different bacterial populations and growth conditions [23; 24], that we also reported in the main text for our experiments on  $E.\ coli$  and  $B.\ subtilis$ . The aim of this final Section is to present theoretical results showing that the property of scale invariance is common to all size distributions, viz.  $s_b$ ,  $s_d$  and  $\Delta_d$ . In other words, if one of the three distributions is scale-invariant, the others as well inherit that property.

The distribution  $\rho_Z(z)$  of a (generic) random variable Z is scale invariant if it has the form:

$$\rho_Z(z) = \frac{1}{\langle Z \rangle} \phi\left(\frac{z}{\langle Z \rangle}\right) , \qquad (36)$$

where  $\phi$  is an arbitrary non-negative normalized function. The statistics of Z is supposed to change with conditions, e.g. of the environment or nutrients. The non-trivial content of the scaling form (36) is that when conditions are varied, the distribution will be modified, yet its shape remains invariant when properly rescaled by the new mean value of Z. The form (36) also implies that, when conditions are varied, the normalized moments  $\langle Z^p \rangle / \langle Z \rangle^p$  will remain constant. Finally, the scaling form (36) is also equivalent to the statement that the Laplace transform  $\mathcal{L}_Z(u)$  of the distribution  $\rho_Z$  has the form  $\mathcal{L}_Z(u) = \psi(u\langle Z \rangle)$ , where u is the Laplace transform variable and  $\psi$  is an arbitrary function (respecting the general constraints for the Laplace transform of a probability distribution).

Let us start by showing that if the scaling form (36) holds for the distribution of the cell size either at division  $s_d$  or at birth  $s_b$ , then it holds also for the other quantity and for the distribution of the added size at division  $\Delta_d$ .

Indeed, if noise in the halving of the sizes at division is neglected,  $s_d = 2s_b$ . The distributions for  $s_d$  and  $s_b$  are then related as  $\rho_b(s_b) = 2\rho_d(2s_b)$  and the scaling form of either one of the distributions clearly implies scale-invariance for the other. Moreover, since  $s_d = s_b + \Delta_d$  and  $\Delta_d$  is independent of  $s_b$ , we have

$$\mathcal{L}_{sd}(u) = \mathcal{L}_{sb}(u) \times \mathcal{L}_{\Delta_d}(u), \tag{37}$$

where  $\mathcal{L}$  indicates the Laplace transform of the respective probability distributions. Using  $s_d = 2s_b$  we have  $\mathcal{L}_{s_b}(2u)/\mathcal{L}_{s_b}(u) = \mathcal{L}_{\Delta_d}(u)$  and therefore the distribution of the added size inherits the scale invariance of  $s_b$  (if the latter has it).

If noise in the halving at division of the size of daughters is included, the argument is slightly more involved. We use  $s_d = \lambda s_b$ , where  $\lambda$  is a random variable centered around 2. We shall assume that the distribution of  $\lambda$  does not change as the means  $\langle s_b \rangle$  and  $\langle s_d \rangle$  vary with growth conditions. Taking the logarithm of  $s_d = \lambda s_b$ , we have again a sum and the Laplace transforms of the logarithms of the three variables are therefore related as in (37). The scale-invariant form (36) implies for the Laplace transform of the distribution of  $\ln Z$  that  $\mathcal{L}_{\ln Z} = \langle Z \rangle^{-u} \psi(u)$ , where  $\psi$  is arbitrary yet it does not contain  $\langle Z \rangle$ . Using that  $\langle s_d \rangle = 2 \langle s_b \rangle = \langle \lambda \rangle \langle s_b \rangle$  (which is valid in any growth condition), one can verify that if either  $s_b$  or  $s_d$  is scale-invariant, the other variable will inherit that property.

Finally, we show that scale invariance (if present) is also inherited by the size distribution  $n(s, \Delta, t)$ . The equation for its dynamics is (13), which reduces to

$$kn(s,\Delta) + \partial_s(g(s)n(s,\Delta)) + \partial_\Delta(g(s)n(s,\Delta)) = -\gamma(\Delta)g(s)n(s,\Delta),$$
(38)

in the stationary state with the growth rate k defined by (16). Taking  $g(s) = \alpha s$  and using  $\langle s \rangle \propto \langle \Delta \rangle$ , one can verify that a scaling form for  $n(s, \Delta)$  is indeed compatible with (38).

A more explicit way to relate  $n(s, \Delta)$  to the distributions of  $\Delta$  and  $s_b$  involves the integration of (38) along the characteristics and the tracking of cells from the current time t back to their last division. For that purpose, it is convenient to introduce the age  $\xi$  of a cell, as in Section III.B, so that

$$\frac{ds}{d\xi} = g(s), \qquad \frac{d\Delta}{d\xi} = g(s), \tag{39}$$

during the elongation of the cell. The initial size  $s_b = s(0)$  and  $\Delta(0) = 0$  are the initial conditions. The equation (38) is rewritten as

$$\frac{dn(s(\xi), \Delta(\xi))}{d\xi} = -F(s(\xi), \Delta(\xi))n(s(\xi), \Delta(\xi)), \qquad (40)$$

where  $F(s, \Delta) = k + \partial_s g(s) + \gamma(\Delta)g(s)$ . We can then track each cell back to its birth:

$$n(s,\Delta) = n(s-\Delta,0)e^{-\int_0^{\xi} d\xi' F(s(\xi'),\Delta(\xi'))} = n(s-\Delta,0)e^{-\int_0^{\Delta} d\Delta' \frac{1}{g(s(\Delta'))}F(s(\Delta'),\Delta')}$$
$$= n(s-\Delta,0)e^{-\int_0^{\Delta} d\Delta' \gamma(\Delta')}e^{-\int_0^{\Delta} d\Delta' \frac{k+\partial_s g(s(\Delta'))}{g(s(\Delta'))}}.$$
 (41)

For the exponential elongation rate  $g(s) = \alpha s \ln 2$ , equation (17) gives  $k = \alpha \ln 2$  and we have that

$$n(s,\Delta) \propto \rho_b(s-\Delta) \left(1 - \int_0^\Delta dx \rho_{\Delta_d}(x)\right) e^{-\int_0^\Delta dx \frac{2}{s(0)+x}} = \rho_b(s-\Delta) \left(1 - \int_0^\Delta dx \rho_{\Delta_d}(x)\right) \left(1 - \frac{\Delta}{s}\right)^2, \tag{42}$$

where we have used (15) to express  $e^{-\int_0^{\Delta} d\Delta' \gamma(\Delta')}$  in terms of  $\rho_{\Delta_d}$ . it is immediate to verify that if  $\rho_b$  and  $\rho_{\Delta_d}$  are scale-invariant, so will  $n(s,\Delta)$  be. Integrating (42) over  $\Delta$ , we obtain for the marginal n(s) at the steady state:

$$n(s) \propto \int_0^s dx \ \rho_b(s-x) \left(1 - \int_0^x dy \rho_{\Delta_d}(y)\right) \left(1 - \frac{x}{s}\right)^2. \tag{43}$$

### VI. FURTHER DETAILS ON THE RELATION BETWEEN FIGURE 1C AND THE CONSTANCY OF THE CELL MASS AT INITIATION OF REPLICATION

W.D. Donachie proposed in [25] that the mass of the cells (per origin of replication) at the time of initiation of the chromosome replication, is independent of the growth rate. Mass and volume will be used interchangeably here as the density is known to be roughly constant. The argument in [25] is based on two premises:

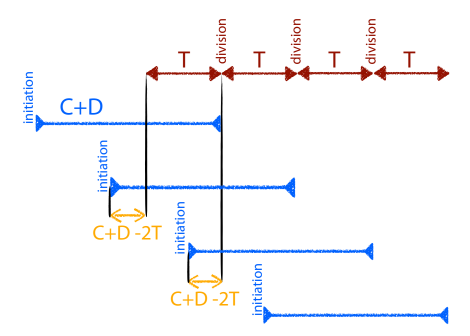


Figure S22. Scheme of the events of initiation of replication and cell division. C denotes the average time for chromosome replication, D is the average time for cell division and the cell average doubling time is denoted here by T. The sum C+D is supposed to be constant for different T. The figure shows that the time separating chromosome initiation to the closest successive cell division is C+D-2T. The scheme drawn here refers to those situations where  $\lfloor \frac{C+D}{T} \rfloor = 2$ , i.e.  $2T \le C+D < 3T$ . In the general case where the integer  $\lfloor \frac{C+D}{T} \rfloor = r$ , i.e.  $rT \le C+D < (r+1)T$ , the corresponding time is C+D-rT.

(i) The sum of the average time C for chromosome replication and the average time D for cell division is constant in diverse growth conditions. Growth conditions are supposed to be fast, i.e. the cell average doubling time, denoted here by T, satisfies  $T \leq C + D$ . The doubling time controls the average volume of cells of age  $\xi$  as

$$\langle V_{\xi} \rangle = \langle V_b \rangle \, 2^{\xi/T} \,, \tag{44}$$

where  $\langle V_b \rangle$  is the average volume at birth. The constancy of C+D was supported by the experiments in Ref. [26] (with  $C+D \simeq 60$  minutes).

(ii) The average volume (mass) of newborn cells scales with T as  $\langle V_b \rangle \propto 2^{\frac{\mu}{T}-1}$ , where  $\mu$  is a constant and T is the doubling time defined above. Such behavior of  $\langle V_b \rangle$  was obtained in Refs. [4; 5] (with  $\mu \simeq 60$  minutes).

Ref. [25] derives the consequence of (i) and (ii) for the time of the initiation of chromosome replication. As shown in Figure S22, the time separating chromosome initiation to the closest successive cell division is C+D-rT, where r is the floor of  $\frac{C+D}{T}$ , i.e. the integer satisfying  $rT \leq C+D < (r+1)T$ . Therefore, the age of cells at the initiation time is T-(C+D-rT)=(r+1)T-(C+D). It follows from (44) and  $\langle V_b \rangle \propto 2^{\frac{\mu}{T}-1}$  that the average volume of cells at the initiation time is

$$\langle V_I \rangle = \langle V_b \rangle \, 2^{r+1 - \frac{C+D}{T}} \propto 2^{r - \frac{C+D-\mu}{T}} \,. \tag{45}$$

The remark in Ref. [25] is that the expression (45) reduces to  $2^r$  if  $\mu = C + D$ , which is precisely the equality obtained in the experiments. Furthermore, it is easy to verify from Figure S22 that the number of origins of replication present at the time of initiation is  $2^r$ . The major ensuing conclusion is that if hypothesis (i) holds and if  $\mu = C + D$ , i.e. if the average volume of newborn cells scales as

$$\langle V_b \rangle \propto 2^{\frac{C+D}{T}-1} \,, \tag{46}$$

then the average mass per origin of replication at the time of initiation is a constant independent of T. Notice that the average volume  $\langle V_b \rangle$  of newborn cells is equal to the average volume  $\langle \Delta V_d \rangle$  added at division, which is the key quantity in the  $\Delta$  model. The average volume added at division is related to the size added at division  $\Delta_d$  discussed in previous sections by the relation  $\Delta V_d = \text{const.} \times w^2 \Delta_d$ , where w is the width of the cell (which stays roughly constant over the cycle of growth) and the constant reflects the rod-like shape of E. coli cells. The equality  $\langle V_b \rangle = \langle \Delta V_d \rangle$  is generally valid since  $\langle V_d \rangle = 2 \langle V_b \rangle = \langle V_b \rangle + \langle \Delta V_d \rangle$ .

Assuming the validity of Donachie's arguments, equation (46) can be used to estimate the value of C+D from the scaling of  $\langle V_b \rangle$  vs T (see, e.g., [21]). This procedure yields the estimation  $C+D \simeq 66$  minutes for E. coli data in Fig. 1C of the main text.

### VII. CONSTANCY OF TOTAL MASS OF CONSTITUTIVE PROTEINS WITH RESPECT TO GROWTH CONDITION

The starting point of Scott et al. [27] is two fundamental correlations between the fraction of ribosomal mass of the proteome,  $\phi_{\rm R}$ , and the average growth rate,  $\lambda$ , for a given growth condition. The two correlations are

$$\langle \phi_{R} \rangle = \langle \phi_{R}^{\min} \rangle + \alpha \langle \lambda \rangle$$

$$\langle \phi_{R} \rangle = \langle \phi_{R}^{\max} \rangle - \beta (\langle \lambda \rangle) \langle \lambda \rangle ,$$

$$(47)$$

$$\langle \phi_{R} \rangle = \langle \phi_{R}^{\text{max}} \rangle - \beta \left( \langle \lambda \rangle \right) \langle \lambda \rangle, \tag{48}$$

where  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.07$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.55$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.49$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.04$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm max}\right>\approx 0.04$  for *E. coli* MG1655 and B/r and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  and  $\left<\phi_{\rm R}^{\rm min}\right>\approx 0.04$  for *E. coli* MG1655 and B/r an NCM3722. The slope  $\alpha = 0.18 \pm 0.1$  is a positive constant for the three different *E. coli* strains. The slope  $\beta$  depends on the growth condition (namely,  $\langle \lambda \rangle$ ), but is constant under a given growth condition. Asymptotically,  $\beta$  can vary between 0 and  $\infty$  as the growth condition changes from rich to poor growth media, respectively.

The above two correlations were obtained from the following experiments. For the first correlation (Eq. 47),  $\langle \phi_{\rm R} \rangle$  of a steady-state population was measured for each growth condition  $\langle \lambda \rangle$ , adding additional information to the growth law (Ref. [4]; Eq. 44; Figure 1C in the main text). For the second correlation (Eq. 48), multiple experiments with varying sub-lethal dosage of chloramphenical (Cam) were performed to measure  $\langle \phi_{\rm R} \rangle$  and  $\langle \lambda' \rangle$  for a given growth condition  $\langle \lambda \rangle$ . Each set of experiments were repeated for other growth conditions.

The invariance of  $\phi_{\rm R}^{\rm max} < 1$  is important. Scott *et al.* [27] interpreted this constancy as that the rest of the proteome fraction,  $1 - \phi_{\rm R}^{\rm max}$ , consists of house-keeping proteins, Q, but in general it can be a basal level of any non-rib second proteins. ribosomal proteins. Importantly, since  $\phi_R^{max}$  is constant,  $\phi_Q = 1 - \phi_R^{max}$  is also constant under all growth conditions. Once we accept this and that the proteome consists of only three types of proteins [ribosomal (R), non-ribosomal (P)and their basal level (Q), we can predict their relative mass fractions for any growth conditions.

Cell cycle proteins and nutrient transporters belong to P sector, and their total mass fraction of the total proteome can be obtained by solving Eqs. 47 and 48 for  $\langle \phi_P \rangle = \langle \phi_R^{\text{max}} \rangle - \langle \phi_R \rangle$ , leading to

$$\langle \phi_{P} \rangle = \frac{\beta \left( \langle \lambda \rangle \right)}{\alpha + \beta \left( \langle \lambda \rangle \right)} \left( \langle \phi_{R}^{max} \rangle - \langle \phi_{R}^{min} \rangle \right) \tag{49}$$

$$= \langle \phi_{\mathbf{R}}^{\mathbf{max}} \rangle - \langle \phi_{\mathbf{R}}^{\mathbf{min}} \rangle - \alpha \lambda. \tag{50}$$

The total mass of the P-sector proteins per cell with respect to the per-cell total proteome can be estimated by multiplying the average volume  $\langle V \rangle$  of the cells for each growth condition and  $\langle \phi_{\rm P} \rangle$ . The growth law is given by  $\langle V \rangle = 0.24 \cdot \langle V_0 \rangle \, e^{0.81 \langle \lambda \rangle}$  for NCM3722 (this work),  $\langle V \rangle = 0.41 \cdot \langle V_0 \rangle \, e^{0.62 \langle \lambda \rangle}$ ,  $\langle V \rangle = 0.33 \cdot \langle V_0 \rangle \, e^{0.70 \langle \lambda \rangle}$ ,  $\langle V \rangle = 0.29 \cdot \langle V_0 \rangle \, e^{0.77 \langle \lambda \rangle}$ , and  $\langle V \rangle = 0.52 \cdot \langle V_0 \rangle \, e^{0.67 \langle \lambda \rangle}$  for B/r H266, B/r K, B/r A, and BilvA thyA, respectively [28].  $\langle \phi_{\rm P} \rangle$ with the experimental parameters  $\alpha, \beta(\langle \lambda \rangle), \phi_{R}^{\min}, \phi_{R}^{\max}$  were reported in Scott *et al.* [27].

For the range of the average growth rate  $\langle \phi_P \rangle$  was measured, the total number of proteins in P sector is remarkably constant in E. coli (Figure 4, main text).

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### VIII. APPENDIX

### A. Background: development of the adder model in this work (by Suckjoon Jun)

The central thesis of our work is that bacterial cells grow by a constant size  $\Delta$  between birth and division. What initially led us to this discovery was the correlations analysis of our published data in [3]. Specifically, we studied the relationship between the newborn cell size  $(s_b)$  and the dividing cell size  $(s_d)$ . Our expectation was that the dividing size would be roughly 2 times the newborn size  $(s_d = 2s_b)$ . However, the linear fit of the correlations systematically deviated from the expected  $s_d = 2s_b$ . Instead, we found  $s_d = s_b + \Delta$  (Figure S23). For almost one year from 2011 to 2012, we repeated our analysis investigating any possible errors in image and data analysis, and always reached the same result,  $s_d = s_b + \Delta$ . In the mean time, we were extending our experiments in [3] to other growth conditions and also other organisms, especially, B. subtilis.

On August 21, 2012, Johan Paulsson visited our lab then at the FAS Center for Systems Biology at Harvard University to discuss cell size control. During our meeting, Taheri-Araghi presented his  $s_{\rm d}=s_{\rm b}+\Delta$ . Upon seeing this, Paulsson made a profound remark that a relationship like " $y=x+\Delta$ " exists in his field of plasmid copy-number control. Importantly, its interpretation is that cells control the average plasmid copy number by adding a constant number  $N_{\rm c}$  of plasmids in each generation, irrespective of the number of plasmids the cell is born with. In the dynamical process, it is straightforward to show that this ensures copy-number homeostasis with the average copy number being  $N_{\rm c}$  itself. In the case of cell size control,  $s_{\rm d}=s_{\rm b}+\Delta$  would then mean that cells sense neither size nor time, but add a constant mass  $\Delta$  irrespective of the initial cell size  $s_{\rm b}$ .

After our meeting with Paulsson, we had to make a decision. On the one hand, we had a potential text-level discovery that virtually no one in bacterial physiology and cell size control were aware of (with one exception of Koppes and his closest colleagues; see Sec. VIII.B). On the other hand, correlations can suffer from degeneracy of different

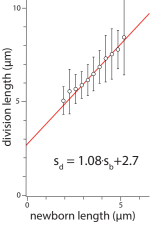


Figure S23.  $s_d = s_b + \Delta$ .

models that result in the same correlation coefficients, mask important causal relationships, but, more important, we only had data for only one growth condition for two  $E.\ coli$  strains (K12 MG1655 and B/r). Due to the limited data, we realized we cannot make a general claim of the constancy of  $\Delta$ . For this reason, we concluded we needed more data from a wide range of physiological conditions as well as different organisms and strains, although it was clear that a complete investigation was going to take several years of major efforts.

Here the experimental difficulty was that *E. coli* changes its dimensions in response to different growth conditions. This is the growth law established in the 1950s, which states that the average size of the cells increases exponentially with respect to the nutrient-imposed growth rate. In practice, what this means is that each growth condition was a project of its own, because each growth condition required a new microfluidic device optimized for the new cell dimensions. By persevering for more than two years, Taheri-Araghi managed to obtain the complete data of quality and quantity that met our standard for *E. coli*. In the mean time, Sauls was extending our experiments to Gram-positive *B. subtilis* and *E. coli* size mutants in collaboration with Taheri-Araghi and Levin's lab.

Starting in late 2012, an intriguing feature emerged from our new data. That is, all probability distribution functions from different growth conditions collapsed when rescaled by their respective means. This is a highly non-trivial result. Fortunately, in early 2013 Vergassola pointed out an important reference by Rinaldo and colleagues [29] on scaling properties of body size, which I consider as important an insight as the one by Paulsson. We immediately started serious collaborations and divided our main responsibilities between experiment and modeling. This Extended Supplementary Information presents the theoretical description by Vergassola and his post-doc Serena Bradde in its final form (see Sec. III, IV, and V), which goes far beyond our preliminary modeling attempt in 2012 (Sec. VIII.C).

To us, obtaining rigorous and relevant experimental data that can pin down a model was the real issue. The idea of constant  $\Delta$  was originally and explicitly proposed by Koppes and colleagues in 1993 [30] and 1997 [31], who called it an "incremental model" (see Sec. VIII.B). In the 1997 article, Koppes wrote

Ideally, what is required is a comprehensive set of data containing distributions of both size and time between successive cellular divisions. Unfortunately, such measurements still have to be made.

This is precisely what we have done in our work.

Finally, size homeostasis requires neither perfect adder nor symmetric division. This explains why different organisms such as yeast or size mutants achieve size homeostasis without having to be perfect symmetric adder. We described this general quantitative adder principle in a recent reference [32].

B. WJ Voorn, LJH Koppes, and NB Grover, Mathematics of cell division in Escherichia coli, Curr Top Mol Genetics 1, 187-194, 1993

To our knowledge, this is the first paper in which Koppes and his colleagues fully developed and presented what they called an *incremental model* (which is identical to our adder principle). For example, they explicitly calculated the correlation coefficients between division and newborn size of  $E.\ coli\ B/r$  strains, and obtained 0.52 and 0.56 for two different growth conditions.

Unfortunately, this reference was published in a journal (Curr Top Mol Genetics) that immediately and completely disappeared right after its first volume from the libraries in the United States as well as online databases. Early this year, one of us (SJ) contacted Voorn, who kept a reprint of this paper and kindly sent a high-resolution picture of every page. For its historical importance, and for the reasons explained above, we include the reference here.

Note that the abstract of this paper reads

Two statistical hypotheses for the occurrence of cell division were tested by observed distributions of cell size during steady-state growth. The thirty year old so-called sloppy-size model could be rejected, whereas the newly-developed incremental-size model was accepted (by lack of alternative).

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Current Topics in Mol. Gen. 1 (1993)

# Mathematics of cell division in *Escherichia coli*: Comparison between sloppy-size and incremental-size kinetics

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1WO statistical hypotheses for the occurrence of cell division "Care tested by observed distributions of cell division "Care tested by observed distributions of cell division "Care tested by observed distributions of cell division care time distribution accurate division of cell-size division accurate of cell-size exist. Instead we have used the most exist.

### TRODUCTION

koch and Schaechter (1962) but forward a model for the statistics of cell division according to which a growing bacterial cell divides when it has attained a critical size; this it has attained a critical size; this size is considered to be the size, on the average, for all the cells in the culture regardless of their size at birth (loch and schaechter 1962). The hypothesis was refined by Powell (1964) and by Hirvey et al. (1967), the latter calculating that it wented to be supported to the size model fits their synchronous growth data rather poorly. The showed that this so-called sloopy-size model fits their synchronous growth data rather poorly. The control of cell division, the incremental-size model, this model states that a growing cell division, the incremental amount; here it is the latter by a critical amount; here it is the latter size that is considered to be the same, on the average, for all the cells in the culture regardless of their size at birth (kabitechek and Woldringh

ideally what is required is a compredeally what is required is a comprehensive date set containing both size and interdivision-time distributions. Unfortunately, no such measurements exist. Instead we have used the most extensive measurements of cell-size distributions available (Dooper 1991, page 657); since the latter were made under the electron microscope on fixed cells, time measurements were of course not possible. Und reil-size measurements were then used to predict various parameters of interdivision-time chosen to be as universal as nossible. The literature was then sear ched for reliable interdivision-time distributions, force we came non the sixteen experiments of for each 1980, and the three additional measurements of the cell-surface areas of dividing cells in both rapid\_ly and slowly growing cultures of sectorishia coll who compute the corresponding interdivision-time distributions for both models under the extension. It appears that the predicted interdivision-time distribution not exponential cell-surface extension, it appears that the predicted interdivision-time distribution in exponential cell-surface dicted interdivision-time distribution is computable with sloppy-size kinetics.

### SCORES ON STATUS

Cultures of s. coli B/r A were grown

to steady state in either nutrient broth (mass doubling time 21 min) or in siyle alannine medium (doubling time 12 min) or time 125 min), fixed in osadum tetra-oxide, and prepared by agar filtration diameter were measured from electron alcregraphs (Grover et al. 1977). Length and cell-surface area computed by assuming an idealized Jeometry of right circular cylinders with hemispherical year used to denote cell-surface area in what follows, size will arrived at in this way.

The relevant equations for the sloppy-smell (1964);

$$y_{k} = 2\sum_{j=0}^{k} (b+j)^{2}/\lambda^{3}$$

t is the interdivision time, q the coefficient of variation, y the skew-ness, the cell size at division, we have ratio of daughter to mother cell size at division, and:

is the parameter of the symmetrical beta distribution of w (hoppes and crown 1992).

It were able to derive analogous expectations for the incremental-size model, but they proved to be intriced the moments of the distribution were calculated actor the assumption of a discrete was all with 47 size classes.

The distribution of incremental size alternative designations for a paper in the literature: life assumption of a discrete was alternative designations for appear in the literature: life assumptions, distribution of a discrete was alternative designations for appear in the literature: life assumption which is a distribution of discrete was alternative designation and division—setj-

increase in size z between birth and division depends not on the size of the newborn cell x nor on its future size at division y but only on the increment y-x. Thus the size distribution of dividing cells  $\phi(y)$  assumes the form:

$$\varphi(y) = 2y' \int_{0}^{y} x \cdot \psi(x) \cdot h(y-x) \cdot dx$$

where w(x) is the size distribution of newborn cells, the factor 2x/y being a consequence of the fact that cell numbers in steady-tate populations increase exponentially (Painter and Marr 1968). This equation complements the one first derived by rowell for the cell-size distribution of newborn cells (hoppes and Grover 1992). You incremental growth in which individual cells increase in size exponentially, then, the distribution of interdivision times is given by:

$$f_1^{-}(\perp) = \ln(2) \cdot 2^{\frac{1}{2}} \int_{-\infty}^{\infty} \mu(x) \cdot h(x)^{\frac{1}{2}} (x - x) dx,$$
 where  $\hat{\tau}$  ( $\ln(2)/\mu$ , the specific growth rate) is the doubling time of the culture.

Pable 1 summarizes the data on interdivision-time distributions found in
the literature. They are the only
published measurements on wildtype
published measurements expopacterial cells in steady-state exponential growth in which the variance
is not excessive (q. 25). It has been

The first three moments of  $f_k(t)$  were calculated and from these the skewness y and four factor  $\beta$  ( $\equiv y/q$ ). Finally,  $\beta$  and y were compared with the exterimental data by means of an exact nonparametric test; the chisquare test of Wilcoxon (Verwer and Hanninga 1980).

Several data sets were used. The first data set 1) is the sixteen measurements reported by Powell (1953). Three of those (designated by Powell as as 4, Cp 1 and Sf 3) show too high a kurtosis (larger than 6), probably due to the presence of anomalous cells in the tails (Koch 1980), and were either dropped (data set 2) or replaced (data set 3) by those from Schaechter et al.

Table 1. Parameters (mean t standard error) of observed interdivision-

189

2144				
0.600 ± 0.069	0.178 ± 0.006	43.2 ± 5.1	16	13
0.746 = 0.098	0.176 ± 0.007	45.2 ± 6.2	13	2
	0.180 ± 0.007	43.1 ± 5.3	16	1
skewness	,co,	(min)		set
-		11 00	n	Data

Table Parameters of calculated interdivision-size(time) distributions

Sample	B	distribution	ion method	mean(units	units)	,o	skewness
æ	769°	φ(y) h(z) h(z)	likelihood moments	2.42	是是是.	0.091	0.673 0.784 1.137
	0	(1)	likelihood moments moments	125.2	HIT HIT HIT	0.128 0.133 0.199	0.410
o	1162	φ(y) h(z) h(z)	noments likelihood moments	16.94 8.56 8.57	是是	0.117	0.559
		£ (7)	likelihood moments moments	21.0	min min min	0.166 0.172 0.246	0.377 0.834 0.0021

precision of division into siblings of about equal size: 5.22 precision of division into siblings of about equal size: 4.35

suggested that complex media give rise to a greater dispersion of interdivision times than do simple media (rowell and Errington 1963), and in an attempt to correct for such an effect, we defined the dimensionless form factor 3 as y/q (Ash and Powell 1962).

From 9x, 9y, and 9x, we were able to be published) to calculate the correlation coefficient r between size at birth and size at division: 0.52 and

0.56 for interdivision times of 21 min and 125 min, respectively. Such high correlation has been reported before (Errington et al. 1955; kopps et al. 1980) and argues against sloppysize control.

Table 2 lists the observed moments of the size distributions at division of two large samples of E. coll B/r A cells growing under steady-state conditions (Grover et al. 1987), together

between incremental-size sloppy-size models

data set	The last	noremental-size	8120	1 lops	1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	u
appent No	Saaple	Probability'	ity	h	Probability.	ty
siemess a	+		+			1
р.	+	+	+		1	1
sk/q a	+		•			1
0'		+	+		,	1

rett the interdivision moments predicted by the two models. As expected from the two models. As expected for all and Errington 1963), q, and q are smaller at the clower crowth rate, remains because the blockedical processes leading to division are more complicated. The average size at division as more complicated. The average size at division as more controlled that the exponential stream could number in steady-stream controlled the pypothesis of Koch and controlled the range of the fight a coefficient of variation for fight about the exponential size and the incremental size of the fight and a reflected in the value of the spronunced different and a scape of the fight and the state of the spronunced different and a scape of the fight and the state of the spronunced different and a scape of the fight and the state of the spronunced different and the shapes of f(t) for both as a second non according to the stage of the stage of

especially as regards skewness. As expected, the form factor is more discrementatory for shapes, because it rejects the incremental-size hypothesis in two instances. The latter finding could be due to both the upset quality of data sets 2 and 3, and the slow growth rate of the bacteria in sample a, in contrast to that in samples 2 and 3, partly invalidating the comparison. We conclude, therefore, that interdivision time is not simply derived from cell size at division, but rather from interdivision size.

Published data. The interdivision-time distribution has been estimated from steady-state liquid culture and found to be possitively skewed (Harvey et al. 1967; warr et al. 1969). This method appears less sensitive than the earlier, direct microscopio observations of individual cells, however, because the cumulative nature if the data can be expected to distort whatever skewness may be present (Harvey 1972; Plank and Harvey 1979).

The earliest distributions of bacterial interdivision time were obtained by Kelly and Rahm (1923) who recorded the family tree of individual cells under the microscope, primarily to estimate mortality. Their skewed distributions show low kurtosis (less than 3) but high coefficient of variation (more than 5%) owing mainly to the low resolution (5 min intervals). This technique was improved considerably by Powell (1955 and 1956) who constructed a special culture chamber that permitted belanced growth under the microscope. The distributions obtained this way are regarded as the most reliable available, and show both low coefficient of variation and low kurtosis (Powell, 1955). Later measurements by Schaechter et al. (1962) and by Kibitschek (1962) confirm the positive skew.

Distributional shape. In contrast to above, we measured the dimensions of dry, dead, flattened bacteria sampled from steady-state cultures, under the electron microscope. Despite this apparent drawback, our cell sizes were found to be approximately proportional to the in vivo cell sizes whereas the precise form of the in vivo interdivision-time distribution remains unknown, its form factor (%) probably lies between 2 and 8, allowing us to reject sloppy-size division control (Koch and Schaechter 1962; probably lies between 2 and 8, allowing us to reject sloppy-size division control (Koch and Schaechter 1962; probably lies between 2 and 8, allowing us to reject sloppy-size division control (Koch and Schaechter 1962; probably lies between 2 and 8, allowing us detail growth implies that many of the old cell-cycle ideas, be they many random events (Rahm 1927; Keniall 1962; Koch 1965; Kubitschek 1971) or more quantitative, and whether they involve one (1984), two (Brooks et al. 1986), are still valled but should be applied to the distribution of interdivision the size or cocept in the hypothesis of the factor (1962) was that size, rather than the f(t). The

ought to be considered the relevant variable (Powell 1984; Errington et al. 1985; Koch 1980). Here we show that it is not absolute size but relative size it.e. size corrected for the size of the cell at birth. Thus, while growth rate is independent of cell age, the division probability (Bell 1868; Kubitschek and Woddringh 1987) is a function of both age and control of cell division presupposes the existence of underlying molecular events in the division pathway, such hypotherical events could occur either with constant probability, exponential kinetics (Tyson 1987), or with a probability that is inversely proportional to cell size, hyperbolic kinetics (Berg and Blowberg 1977), depending on genotype (Gottesman 1939) in either case, the resulting interdivision—time distribution will not be symmetrical (Painter and Marr 1967) but Yule-like.

reasons for the survivel of Koch and Schaechter's sloppy-size model was its ability to predict with reasonable accuracy the correlation between the intendivision time of parents and that of their progeny, -0.45, and between the siblings themselves, +0.35 (Koch and Schaechter 1952; Fowell and Errington 1963; Kubitschek 1960). The correlations predicted by the incremental-size model are lower, -0.20 and +0.20, respectively (unpublished results). A closer look at the experimental data, however, suggests that discrimination between these two sets of values is not at all reliable:
microscopic observation artificially random errors in the rejistry of cell separation (Fainter 1975) while Gaussian filtering of synchronous-growth curves tends to do the same owing to the inherent assumption of symmetry (Plank and darrey 1979). It has receively been demonstrated quite convincingly, by using a chromosomal runcaway replication mutant, that the UNAreplication and the cell-division of the same separate consequences of the same spare intendependent, i.e. they are separate consequences of the same symmetry (1975) and the convincingly of unpublication division of the same start that the UNAreplication and the reliables of the same symmetry of the same spare intendependent, i.e. they are separate consequences of the same symmetry (1980).

Table 1

Able 2

Probability that difference between predicted and experimental probability that difference between predicted and experimental probability that difference between the chi-square test for symmetry (Verwer and Kunninga, 1980). On the assumption of symmetry (Verwer and tenninga, 1980), on the assumption of equivalence of both methods, i.e. moments and maximum likelinoid (see Appendix), the average predicted parameter (Table 2) mas tested by observed values (Table 1) at x = 0.005 significancy.

are timed by the same of the cell are timed by the same incrementalsize mechanism as that proposed here for cell division. Sibling correlation them be expected to contribute to the tight synchrony of initiation obserted to occur at multiple origins in the same cell (Koppes et al. 1978; Growth law, In 1986), (Koppes 1992) and of the

Crover et al. 1987), we compared sight growth andels for surface extension during unrestricted balanced strowth two exponential and six rejected on statistical grounds, the fit to the experimental data being imadequate. For theoretical grounds, the imadequate. For theoretical grounds, the imadequate for theoretical grounds, the rate doubles by a sloppy-size mechanism. There are thus four surviving one-parameter models, two bilinear and two exponential ones. It may be that individual bacterial cells in continuous cultures grow bilinearly with discrete rate doublings at muclear division (Kubitschek 1986; fubitschek and Pai 1983), but the situation in batch cultures remains equivocal. We chose simple exponential extension of the cell-surface of couptial and not the extra parameter of being justified by the data; the introduction of the extra parameter in the data; the arguments presented here interests is very small (Crover et al. 1987) and not expected to detract in a strong the skenness of the inter-

The property of the state of their cell cycles are street (Cooper 1991). It is temptered (Cooper 1991). It is temptered (Cooper 1991). It is temptered to consider extending a state of incremental este such as a state of the st

Synchronous growth. The rejection of the sloppy-size mechanism has implications for the way in which gellope outless for the way in which gellope over the specific blocked a reactions should be investigated. Apart from planting possible trauma caused by the synchronization procedure itself, selection by age (membrane elution) is as poortells by size (centrifugal elutriacells by size (centrifugal elutriacells by size (centrifugal elutriacells) as far as the cell cycle is concerned. It would appear that the only reliable method is double selection, by both age and size (No pee et al. 1980).

bution of the size ratio w-sibling/parent is assumed to be B(b;w), where be denotes the single parameter of the beta division imprecision distribution (Koppes and Grover 1992). With then straightforward to calculate from given function h(z) the birth and division functions w(x) and w(y), respectively. Sufficient accuracy was obtained by taking as unit of cell size about one tenth of the straidard deviation of the observed distribution of the discrete division probabilities h(z) that were tried to maximize the likelinood of the observed parental size frequencies, were allowed to have and were only restricted to be unimodal. Several starting points in a confict the same point of convergence was found.

Of h(z)-probabilities were tried, but every time the same point of convergence was found. It is then possible to calculate the moments & of the distribution of state at division. It is then possible to calculate the moments & of the distribution size z from

$$\bar{y}_{k} = 2 \sum_{j=0}^{K!} {\binom{k-1}{j}} \cdot \bar{x}_{k j} \cdot \bar{z}_{j}, k = 1, 2, ...$$

(An elementary proof is by Laplace transformation of the incremental-size equation, derivation and substitution of zero argument). From Powell's equation it follows that:

where b, = }

bk .

1,

Let g(t) denote the distribution interdivision—size ratio t=z/x exp( $\mu\tau$ ) -1. Then:  $b_k = b_{k-1}(b+k-1)/(2b+k-1)$ H 11 0%

B(t) =  $\int_{-\infty}^{\infty} \psi(x).$ 

wellin transformation gives the moments  $t_k$  as product of those and  $\phi(y)$ : of

where bo = X.x . Zx 11 p\_K . Y.K . K II

Finally, the approximate moments  $\tau_k$  are found by linear transformation of g(t):  $\tau_k$   $\tau_k^2 \ln(2) + 1 - 1/2 \ln(2)$ . This transformation only affects the mean and standard deviation, but not the coefficients of skewness and kurtosis. b<sub>1-K</sub>·(2b-k)/(b-k) k = 2,

We wish to express our thanks to dr. Nanninga and to dr. Woldringh for their kindness and warm hospitality, respectively.
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C. Preliminary theory note by Taheri-Araghi, September 28, 2012

For readers interested in the initial development on the theory front, we included an original preliminary theory note written by Taheri-Araghi. This summarizes our understanding of the consequences of constancy of  $\Delta$  as of September 2012 soon after our meeting with Johan Paulsson. This was primarily at the correlations level based on limited date sets in Wang  $et\ al.\ [3]$ . As explained in Section VIII.A, a complete, independent theoretical framework was established only after Vergassola and Bradde joined with the on-going acquisition of new experimental data.

### Growth and Cell Size Control in ${\it Escherichia~coli}$

Sattar Taheri-Araghi, Ariel Amir, Suckjoon Jun<br/> September 28, 2012

### 0.1 Introduction

The life cycle of an *Eschirichia coli* cell from an outsiders point of view consist of growth in size, ended by division of the cell in half. Complex molecular mechanisms drive and control this cell cycle, on which our knowledge is still at early stages. A precise quantitative picture of cell size evolution over a cell cycle, and coordination between size and division can shape our fundamental understanding on growth, cell cycle, and the molecular circuits that control this process under the hood.

In this study, we investigate the coordination between cell size and division of *E. coli* during steady state growth. Specifically, we challenge the general belief that cells "double up" their size during their life cycle. We analyzed growth of thousands of *E. coli* cells in "mother machine", a high throughput continues culture microfluidic device.

Quantitative analysis of cell level parameters, such as size, elongation rate and generation time, reveals a remarkable, yet simple, coordination between division and cell size: the cell mass increase during each cell cycle is constant regardless of the cell size at birth. That is, *E. coli* cells somehow measure the amount of mass they accumulate after birth. Once reaching a critical value, depending on the growth condition, cells divide into two viable daughters. This finding argues against models that propose cells measure their size to trigger division and models that picture a constant timer controlling cell division. However, at a population average, it is consistent with that the cell size at division is twice the size at birth.

### 0.2 Steady state growth from a single cell perspective

We used mother machine to collect data on the growth and cell cycle of 7,300 single cells in the steady state growth. Images of growing cell, collected by time lapse microscopy, analyzed by custom developed image analysis software and single cell parameters are quantified essentially in terms of four independently measured parameters:

- $\ell$ : the length of a cell at birth (newborn length)
- L: the length of a cell just before division (division length)
- T: the generation time of a cell
- $\bullet$   $\lambda$ : the elongation rate, as defined for one dimensional growth of a cell below.

In this work we, put an emphasis on precise quantitative analysis of the data obtained from the experiment. Specifically, we investigate not only the average and fluctuations of different parameter, but also the correlation among them and eventually develop an analytical model to explain the results and provide insights on cell size control mechanism in *E. coli* cells.

### 0.2.1 How big is the variations in cell size?

A previous study has shown  $E.\ coli$  cells have a robust growth mechanism over hundreds of generation where the their elongation rate remain constant regardless of the replication age of cell. In other words, cells do not get slow in growth as they age, even though the probability of fillamentation becomes larger. While cells maintain a constant elongation rate and robust growth, their size and generation time do fluctuate over 20% of the average value. This level of size variations indicates that there should be a mechanism that controls the fluctuations in a populations of the cells. An obvious question in this context is that what happens to those cells who are much larger, or smaller, than the average population size? How do they catch up with the average size?

Fig. 1 shows histograms of cell size at birth (left panel) and division length (right panel). The distributions are not symmetric, later in this study we show that they are approximately lon-normal distributions.

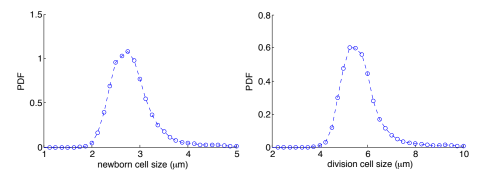


Figure 1: Normalized distribution of newborn cell length (left panel) and cell length at division right panel. Distributions are asymmetric (non Gaussian).

Fig. 2 demonstrates an scatter plot of division length as a function of newborn length. The red line that passes through the scatter points is the best line fit to this data points. There are two points in this graph that deserve some discussion. First, the division length is correlated with the newborn length. This implies that, if a cell is born larger than the average, its size at division will be likely larger than average too, yielding larger than average daughter cells. While this may seem inconsistent with an active mechanism controlling the cell size, the second thing we want to point out is the slope of the fitted line. Consider, hypothetically, the slope of the line was two. That would imply the size of a cell at division is twice the newborn size resulting in daughter cells with same size as previous generation. A slope smaller than two, however, indicates that daughter cells have a size closer to the average size compared to the newborn cells of the previous generation.

What is evidence here is that, cells who deviate from average size do get back to the average size but this does not happen in one generation. In the next section, we show how cells coordinate growth to catch up with an average size of the cell.

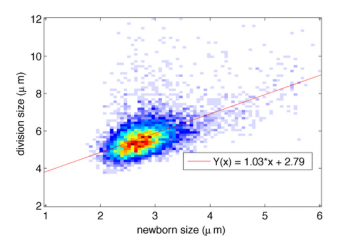


Figure 2: Length of cells right before division as a function of their newborn length for a total number of 7,300 mother cells in steady state growth. The slope of the fitted line, 1.03, is an indication of an active mechanism controlling cell size as opposed to *doubling* cell mass during a cycle.

### 0.2.2 How do cells control their size?

At a cell level observation, we can quantify the growth in terms of elongation rate and generation time. Elongation rate denotes how *fast* cells grow and generation time denotes *how long* they grow before division. Here, we illustrated the correlation of these two parameters with newborn cell size.

Figure 3 depicts scatter plots of cell elongation rate (left panel) and generation time (right panel) versus newborn cell size. Elongation rate does not show any significant correlation with cell size. On the other hand, generation time shows a strong negative correlation with newborn size.

To further analyze this observation, we have binned cells based on their newborn size and plotted the average elongation rate and average generation time in each bin. Figure 4 shows the average elongation rate (left panel) and average generation time (right panel) of each bin as a function of newborn cell size. The error bars show the standard deviation in each bin. For both elongation rate and generation time, the standard deviations seems similar in different bins.

One important observation here is that normalized standard deviation for generation time is larger than that of elongation rate. For a quantitative com-

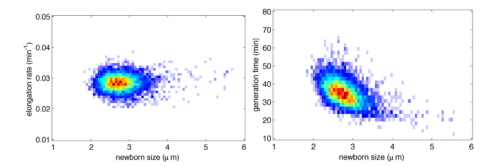


Figure 3: Scatter plot of elongation rate (left panel) and generation time (right panel) as a function of newborn cell size. Elongation rate does not show any correlation with size while cells generation time is negatively correlation with their size at birth.

parison, we have chosen one bin and normalized a histogram of both variables as plotted in figure 5. The generation time has a standard variation of  $\sigma = 0.144$  while the elongation rate standard deviation is sigma = 0.077. In such stochastic systems, the parameter with a higher fluctuation eventually dominates the systems fluctuation. In the next section, we use this fact to reduce the complexity of our theoretical model considering all the fluctuations in size come from variations in generation times.

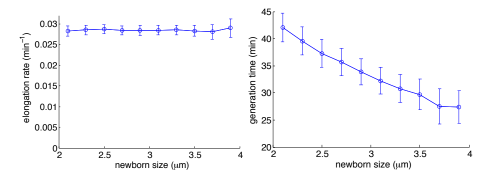


Figure 4: Elongation rate and generation time as a function of newborn cell size.

### 0.2.3 Experiment condition

The experiment is conducted on SJ288 strain in rich MOPS media at 30C. Mother machine device was made from SJ110 microchannel set. The growth media was continuously injected to the device at 3,000  $\mu$ l/hr. Microscopy is performed with Nikon Ti using 100X oil immersion lens with bright field phase contrast. Images from the mother machine device acquired at 10 different fields

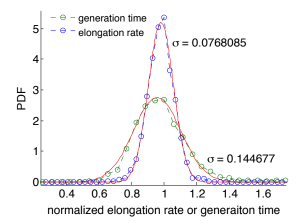


Figure 5: Normalized elongation rate and generation histogram for a bin of figure 4. The generation time has a wider distribution with  $\sigma = 0.145$  compared to elongation rate with  $\sigma = 0.077$ .

of views at 2 min intervals. LED illumination was used at 100% power with exposure time of 70ms for each image acquisition.

### 0.3 Quantitative model for growth

### 0.3.1 Quantitative model for in depth understanding of cell size control mechanism

Based on our experimental observations, we put forward an analytical model for growth and cell cycle of  $E.\ coli$  cells. Our model essentially describes the strategy that  $E.\ coli$  cells apply to control their growth and division in each cell cycle. The model is at a single cell level, yet has the power to predict the distributions and correlations between cell size, elongation rate and generation time. In the following, we present the model and then test the model with the data we have collected from  $E.\ coli$  cells.

Assumptions of the model. The model is constructed upon a few assumptions that we have itemized below. This assumptions are based on our single cell observation.

- Growth of *E. coli* in steady state is one dimensional. That is, the cell maintains a constant width during the growth and upon division. The mass of the cell is thus proportional to the length of the cell. Here and in what follows, length and elongation rate is used to characterize "size" and "size increase rate" of the cells.
- Cells are evidenced to grow exponentially during a cell cycle. Quantitatively, the relationship is  $L = \ell \times 2^{\lambda T}$  where L the is cell length at division,

 $\ell$  the newborn length,  $\lambda$  the elongation rate, and T is the generation time.

• Elongation rate is constant while generation time is a stochastic variable correlated with cell size. As demonstrated in the previous section, the coefficient of variations of elongation rate is two folds smaller than that of generation time. That is, it's effect on cell size variation is about 4 times smaller than that of generation time. To establish a robust model with the smallest number of fitting parameters, we assume the elongation rate, λ, is constant during a cell cycle and is the same for all the cells in the population. On the other hand, the generation time, T, is a stochastic variable, correlated with newborn cell size. This correlation and the noise and fluctuations of T will be discussed below.

Functional form of generation time. The generation time can be divided into two compartments: the deterministic part and the noise part,  $T = T_n + T_d$ . The deterministic part is a function of the newborn cell size, yielding the correlation between generation time and cell size,  $T_d = T_d(\ell)$ . The noise part,  $T_n$ , consistent with the observations, is assumed to be a Gaussian variable with a constant coefficient of variations, independent of the cell size. As the first step for developing the model, we work out the functional form of the deterministic part of the generation time,  $T_d$ . To keep the generality of the model, we construct the function such that it interpolates between two plausible cell cycle models, constant timer model and constant division size model (see below for details). We, however, argue that non of these limiting models are correct and we conclude that cells behavior is somewhat in between, which is explained quantitatively as a model in which cell mass increase is the same in each cell cycle, regardless of the newborn size. This model predicts that any deviation from average cell size will be corrected in a few generations and cell size fluctuations remain constant over different replicative age. An interpolating parameter, which is eventually determined by fitting, indicates in which model the cells are behaving.

Asymptotic models. The two asymptotic models for cell cycle that we interpolate between are constant generation time and constant cell size at division. In the former, cell adjust the generation time to the average expected value depending on the growth condition regardless of the newborn cell size. In the latter, a cell grows until it reaches a especial size, let's say  $2 \times \bar{\ell}$ , before division. Generation time for these limiting cases can be formulated, respectively, as:

$$T_d = \frac{1}{\lambda} \tag{1}$$

$$T_d = \frac{1}{\lambda} + \frac{1}{\lambda} \log \left( \frac{\overline{\ell}}{\ell} \right) \tag{2}$$

Interpolation function. These two limiting models may sound appealing, but we can readily argue that non of these may hold for *E. coli* growth; in the constant timer model, the fluctuations in size only arises from noise in the generation time. In such a model, noise in the cell size accumulates over

generations – no mechanism to control the fluctuations. In the constant division size model, cells divide upon reaching an especial cell size appropriate for cell division. in this model, there will be no correlation between cell size at division and newborn cell size. We use an interpolating parameter  $\alpha$  to bridge between these two cases. The generation time is then defined as

$$T_d = \frac{1}{\lambda} + \frac{\alpha}{\lambda} \log \left( \frac{\overline{\ell}}{\ell} \right) \tag{3}$$

The interpolation parameter  $\alpha$  can be eventually obtained by fitting Eq. 3 to experimental results. The  $\alpha$  obtained from fitting indicates the cells behavior in growth and cell size control.

Special case at  $\alpha=0.5$ : constant mass increase in each cycle. Before proceeding to fitting the generation time to experimental data. We would like to point out that the mid point of the interpolation, at  $\alpha=0.5$ , is an special case that the length increase in each cell cycle is constant  $\bar{\ell}$ . This can be shown as below:

$$L = \ell \times 2^{\lambda T_d}$$

$$\bar{\ell} = \Delta L = \ell \times 2^{\lambda T_d} - \ell$$

$$\bar{\ell}/\ell + 1 = 2^{\lambda T_d}$$

$$\log (1 + \bar{\ell}/\ell) = \lambda T_d.$$
(4)

since  $\bar{\ell}/\ell \approx 1$ , at the first order approximation,  $\log (1 + \bar{\ell}/\ell) \approx 1 + 0.5 \log (\bar{\ell}/\ell)$ . Substituting it back in Eq. 4, we obtain:

$$T_d = \frac{1}{\lambda} + \frac{0.5}{\lambda} \log \left( \frac{\overline{\ell}}{\ell} \right) \tag{5}$$

comparing Eq. 5 with Eq. 3, we see  $\alpha=0.5$  approximately corresponds to the case that cell length increase is constant,  $\bar{\ell}$ , during each cell cycle, regardless of the newborn size.

Derivation of the distributions. Here we derive an analytical expression for the distributions of newborn cell size and generation time. Starting from the generalized form of  $T_d$  in Eq. 3 we can proceed calculating newborn size of a new generation based on the previous generation parameters, assuming precise cells division.

The generation time is given as,

$$T = T_d + T_n$$

$$T = \frac{1}{\lambda} + \frac{\alpha}{\lambda} \log\left(\frac{\overline{\ell}}{\ell}\right) + T_n.$$
(6)

Length of a new generation can be written in terms of previous generation parameters:

$$L = \ell \times 2^{\lambda T}$$

$$\ell_{new} = \frac{1}{2} \ell_{prev} \times 2^{\lambda T}. \tag{7}$$

Substituting Eq. 6 in Eq. 7, and reordering the results we obtain:

$$\log\left(\frac{\ell_{new}}{\overline{\ell}}\right) = \log\left(\frac{\ell_{prev}}{\overline{\ell}}\right) + \lambda T_d + \lambda T_n - 1$$

$$= \log\left(\frac{\ell_{prev}}{\overline{\ell}}\right) + \alpha \log\left(\frac{\overline{\ell}}{\ell_{prev}}\right) + \lambda T_n$$

$$= (1 - \alpha) \log\left(\frac{\ell_{prev}}{\overline{\ell}}\right) + \lambda T_n$$
(8)

From stationarity,  $P\left[\log\left(\ell_{new}/\overline{\ell}\right)\right] = P\left[\log\left(\ell_{prev}/\overline{\ell}\right)\right] = P\left[\log\left(\ell/\overline{\ell}\right)\right]$ . Since the  $T_n$  is a Gaussian variable,  $P\left[\log\left(\ell/\overline{\ell}\right)\right]$  is Gaussian too. Variance of the distribution can be related as follows:

$$\operatorname{Var}\left[\log\left(\ell/\overline{\ell}\right)\right] = (1-\alpha)^{2} \operatorname{Var}\left[\log\left(\ell/\overline{\ell}\right)\right] + \lambda^{2} \operatorname{Var}\left[T_{n}\right],$$
  

$$\alpha(2-\alpha) \operatorname{Var}\left[\log\left(\ell/\overline{\ell}\right)\right] = \lambda^{2} \operatorname{Var}\left[T_{n}\right]$$

$$\operatorname{Var}\left[\log\left(\ell/\overline{\ell}\right)\right] = \frac{\lambda^2 \operatorname{Var}\left[T_n\right]}{\alpha(2-\alpha)} \equiv \sigma_{\ell}^2. \tag{9}$$

Assuming

$$P[T_n] = \frac{1}{\sigma_n \sqrt{2\pi}} \exp\left(\frac{-T_n^2}{2\sigma_n^2}\right),\tag{10}$$

with  $\sigma_n^2 = \text{Var}[T_n]$ , distribution of  $\ell$ , the newborn cell size, can be derived as follows:

$$P\left[\log\left(\ell/\overline{\ell}\right)\right] = \frac{1}{\sigma_{\ell}\sqrt{2\pi}} \exp\left(\frac{-\log\left(\ell/\overline{\ell}\right)^{2}}{2\sigma_{\ell}^{2}}\right)$$

$$P\left(\ell\right) = \frac{1}{\ell\sigma_{\ell}\sqrt{2\pi}} \exp\left(\frac{-\log\left(\ell/\overline{\ell}\right)^{2}}{2\sigma_{\ell}^{2}}\right), \tag{11}$$

with  $\sigma_{\ell}^2 = \lambda^2 \sigma_T^2 / \alpha (2 - \alpha)$ . (see Eq. 9).

The distribution of the generation time can also be calculated in a similar way. Recalling Eq. 6, considering the Gaussian distribution of  $T_n$  and  $\log(\ell/\bar{\ell})$ , the generation time, T, should be a Gaussian variable. The variance is determined from Eq. 6:

$$\operatorname{Var}\left[T\right] = \left(\frac{\alpha}{\lambda}\right)^{2} \operatorname{Var}\left[\log\left(\frac{\overline{\ell}}{\ell}\right)\right] + \operatorname{Var}\left[T_{n}\right],$$

$$\operatorname{Var}\left[T\right] = \frac{2\alpha}{\alpha(2-\alpha)} \operatorname{Var}\left[T_{n}\right] \equiv \sigma_{T}^{2}.$$
(12)

The distribution of generation time, T, is thus given,

$$P[T] = \frac{1}{\sigma_T \sqrt{2\pi}} \exp\left[-\frac{(T - 1/\lambda)^2}{2\sigma_t^2}\right]. \tag{13}$$

### 0.3.2 Testing the model

In this section we test, the model that we have developed against our experimental data. Here are the steps we take to test the mode: i) finding  $\alpha$  by fitting the Eq. 3 to the average generation time versus newborn cell size (Fig. 4, right panel). ii) Finding the variance of  $T_n$  from results plotted in figure 5. iii) comparing the distribution of generation time and cell size, prediction of the model in Eqs. 13 and 11.

Finding the value of interpolating parameter  $\alpha$ . Figure 6 shows the deterministic part of the generation time as a function of newborn cells size for a few different cases including the best fit for  $\alpha$ . The green dashed line in this figure corresponds to constant timer where  $T_d = 1/\lambda$ , regardless of the cell size. The green dashed line corresponds to fixed division size where  $T_d = 1/\lambda + (1/\lambda)\log\left(\overline{\ell}/\ell\right)$ , with  $\overline{\ell}$  the half of the division size. The blue line in the figure with error bars are experimental results and the solid red line is the best fir of Eq. 3 to the experimental results.

The value of  $\alpha$  we have obtained from this fitting is  $\alpha = 0.48$  which is very close to the special case  $\alpha = 0.5$  in which the cell size increase in each cell cycle is constant.

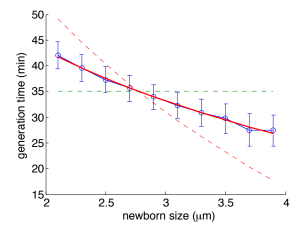


Figure 6: The deterministic part of generation time as a function of newborn size. This graph to to show how to finding the value for the interpolating parameter  $\alpha$ . The green dashed line depicts constant timer model,  $\alpha=0$ . The red dashed line depicts constant division size,  $\alpha$ . The blue line show the experimental results. The solid red line shows the best fit corresponding to  $\alpha=0.48$ .

Finding the standard variation of  $T_n$ . To find the standard deviation of  $T_n$ , we used data similar to that plotted in figure 5 for different bins and took the average. The value we found is  $\sigma_n = 5.38 \,\mathrm{min}$ .

comparing the model versus experiment. Now that we have the two parameters that we had to obtain by fitting,  $\alpha = 0.48$  and  $\sigma_n = 5.38$  min, we can compare the distribution of newborn cell length and generation time as predicted in Eqs. 11 and 13, respectively, with experimental results. Figure 7 depicts both theoretical predictions and experimental results for these distributions. The agreement between theory and experiment is remarkable.

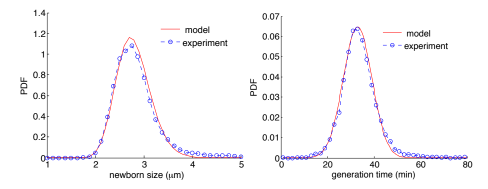


Figure 7: Comparison of model predictions (solid red lines) with experimental data for the distributions of newborn cell size (left panel) and generation time (right panel).

### 0.3.3 Autocorrelation functions

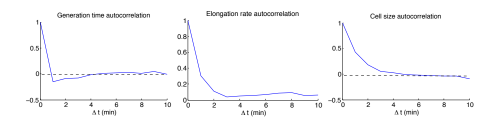


Figure 8: Autocorrelation function for generation time, elongation rate and cel size.

### 0.3.4 Analytical solution for generation time autocorrelation

Here we find the autocorrelation function of generation time for  $\alpha = 0.5$ , as it corresponds to constant cell mass (or, alternatively, length) increase per cell

cycle. The normalized autocorrelation function is defined as

$$A(t) = \frac{\left\langle (T_i - \overline{T}) \cdot (T_{i+t} - \overline{T}) \right\rangle}{\left\langle (T_i - \overline{T})^2 \right\rangle}$$
(14)

where  $T_i$  is the generation time of a cell with replicative age of i and  $\overline{T} = 1/\lambda$  is the average generation time of all cells.  $T_{i+t}$  denotes the generation time of the cell born at replicative age of i+t in the same lineage as the cell with  $T_i$  generation time. The average  $\langle \rangle$  runs over all cells with all replicative ages. In this section we calculate the A(1) and A(2).

To begin with, let's choose a random cell, whose replicative age and newborn length are respectively i and  $\ell_i$ .  $\ell_i$  is a stochastic variable with log-normal distribution as calculated in Eq. 11. The generation time for this cell,  $T_i$ , is given based on Eq. 6:

$$T_i = \frac{1}{\lambda} + \frac{1}{2\lambda} \log\left(\frac{\overline{\ell}}{\ell_i}\right) + T_n^i. \tag{15}$$

where  $T_n^i$  refers to the noise in generation time for this cell. Note that  $T_n^i$  is a Gaussian variable independent of i and  $\ell_i$ . To calculate A(1), we need to go one step forward and find the generation time for the next generation cell, whose replicative age is i + 1. First, considering constant mass increase in each cell cycle, the newborn length for the next generation cell is given by

$$\ell_{i+1} = \frac{1}{2} \left( \overline{\ell} + \ell_i \right) \times 2^{\lambda T_n^i} \tag{16}$$

Then, the generation time corresponding to this newborn length is

$$T_{i+1} = \frac{1}{\lambda} + \frac{1}{2\lambda} \log \left( \frac{2\overline{\ell}}{\ell_i + \overline{\ell}} \times 2^{-\lambda T_n^i} \right) + T_n^{i+1}$$
$$= \frac{1}{\lambda} - \frac{1}{2\lambda} \log \left( \frac{1}{2} + \frac{\ell_i}{2\overline{\ell}} \right) - \frac{1}{2} T_n^i + T_n^{i+1}. \tag{17}$$

Since  $\ell_i/\bar{\ell} \approx 1$ , at the first order approximation we can replace  $\log (1/2 + \ell_i/2\bar{\ell})$  with  $\frac{1}{2} \log (\ell_i/\bar{\ell})$ .  $T_{i+1}$  then reads

$$T_{i+1} = \frac{1}{\lambda} - \frac{1}{4\lambda} \log \left( \frac{\ell_i}{\overline{\ell}} \right) - \frac{1}{2} T_n^i + T_n^{i+1}. \tag{18}$$

Substituting Eqs. 15 and 18 in autocorrelation function we obtain

$$A(1) = \frac{\left\langle \left(\frac{1}{2\lambda}\log\left(\overline{\ell}/\ell_i\right) + T_n^i\right) \cdot \left(-\frac{1}{4\lambda}\log\left(\ell_i/\overline{\ell}\right) - \frac{1}{2}T_n^i + T_n^{i+1}\right)\right\rangle}{\left\langle \left(\frac{1}{2\lambda}\log\left(\overline{\ell}/\ell_i\right) + T_n^i\right)^2\right\rangle}.$$
 (19)

Here the averages are on over all possible  $\ell_i$ ,  $T_n^i$ ,  $T_n^{i+1}$ . Mathematically,  $\langle X \rangle = \int X P\left[\ell_i\right] P\left[T_n^i\right] P\left[T_n^{i+1}\right] d\ell_i dT_n^i dT_n^{i+1}$ . Since  $\log(\ell_i/\bar{\ell})$ ,  $T_n^i$ ,  $T_n^{i+1}$  are independent stochastic variables with Gaussian distribution around zero, many terms

in the above equation are zero once the averaging is carried out. With some simplification we obtain

$$A(1) = \frac{\left\langle \frac{1}{8\lambda^2} \log^2\left(\ell_i/\overline{\ell}\right) - \frac{1}{2} \left(T_n^i\right)^2 \right\rangle}{\left\langle \frac{1}{4\lambda^2} \log^2\left(\overline{\ell}/\ell_i\right) + \left(T_n^i\right)^2 \right\rangle} = \frac{\frac{1}{8\lambda^2} \left\langle \log^2\left(\ell_i/\overline{\ell}\right) \right\rangle - \frac{1}{2} \left\langle \left(T_n^i\right)^2 \right\rangle}{\frac{1}{4\lambda^2} \left\langle \log^2\left(\overline{\ell}/\ell_i\right) \right\rangle + \left\langle \left(T_n^i\right)^2 \right\rangle}$$
(20)

The second moment of the Gaussian variables are given as

$$\left\langle \left(T_n^i\right)^2\right\rangle = \sigma_n^2 \tag{21}$$

$$\langle \log^2 \left( \ell_i / \bar{\ell} \right) \rangle = \sigma_\ell^2 = \frac{\lambda^2 \sigma_n^2}{\frac{1}{2} (2 - \frac{1}{2})}$$
 (22)

Substituting these moments in previous equation we get

$$A(1) = \frac{\frac{1}{4(2-1/2)}\sigma_n^2 - \frac{1}{2}\sigma_n^2}{\frac{1}{2(2-1/2)}\sigma_n^2 + \sigma_n^2} = -\frac{1}{4}.$$
 (23)

To find A(2) we should go one step forward. A similar approach yields

$$A(2) = -\frac{1}{8} \tag{24}$$

### 0.4 conclusions